


# Roadmap on Atomtronics: State of the art and perspective <sup>F</sup>

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 L. Amico,  M. Boshier, G. Birkel, et al.

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## ABSTRACT

Atomtronics deals with matter-wave circuits of ultracold atoms manipulated through magnetic or laser-generated guides with different shapes and intensities. In this way, new types of quantum networks can be constructed in which coherent fluids are controlled with the know-how developed in the atomic and molecular physics community. In particular, quantum devices with enhanced precision, control, and flexibility of their operating conditions can be accessed. Concomitantly, new quantum simulators and emulators harnessing on the coherent current flows can also be developed. Here, the authors survey the landscape of atomtronics-enabled quantum technology and draw a roadmap for the field in the near future. The authors review some of the latest progress achieved in matter-wave circuits' design and atom-chips. Atomtronic networks are deployed as promising platforms for probing many-body physics with a new angle and a new twist. The latter can be done at the level of both equilibrium and nonequilibrium situations. Numerous relevant problems in mesoscopic physics, such as persistent currents and quantum transport in circuits of fermionic or bosonic atoms, are studied through a new lens. The authors summarize some of the atomtronics quantum devices and sensors. Finally, the authors discuss alkali-earth and Rydberg atoms as potential platforms for the realization of atomtronic circuits with special features.

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## TABLE OF CONTENTS

|  |   |  |    |
|--|---|--|----|
| I. INTRODUCTION .....                                | 3 | D. Hybrid atomic-superconducting quantum systems .....                   | 7  |
| II. DYNAMICALLY SCULPTED LIGHT .....                 | 4 | E. Concluding remarks and outlook .....                                  | 7  |
| A. Fast-scanning AODs .....                          | 4 | III. IMPLEMENTING RING CONDENSATES .....                                 | 8  |
| 1. Feed-forward control .....                        | 4 | A. General features of ring traps .....                                  | 8  |
| 2. Phase evolution in time-averaged potentials ..    | 5 | B. Techniques based on magnetic traps .....                              | 8  |
| 3. Atomtronics with time-averaged optical traps.     | 5 | 1. RF dressing and bubbles .....   | 9  |
| B. Optical potentials with liquid-crystal SLMs ..... | 5 | 2. Waveguides formed from time-averaged adiabatic potential (TAAP) ..... | 9  |
| C. Direct imaged DMD optical potentials .....        | 6 | 3. Dynamical ring in an rf-dressed adiabatic bubble potential .....      | 11 |
| 1. Half-toning and time-averaging .....              | 6 | C. Trapping in rings with optical potentials .....                       | 12 |
| 2. Atomtronics with DMDs .....                       | 6 |  |    |
| 3. Turbulence with DMDs .....                        | 6 |  |    |

|  |    |   |    |
|--|----|---|----|
| 1. Optical trapping .....                            | 12 | 2. Spectrum .....                                   | 44 |
| 2. Optical ring traps .....                          | 12 | 3. Metastability .....                              | 45 |
| 3. Imperfections in optical traps .....              | 15 | D. Thermal and quantum phase slips                  |    |
| D. Hybrid traps: RF bubble plus light sheet(s) ..... | 15 | in a one-dimensional Bose gas on a ring .....       | 47 |
| E. Concluding remarks and outlook .....              | 16 | 1. Model and methods .....                          | 47 |
| IV. ATOMTRONIC CHIPS AND HYBRID SYSTEMS .....        | 16 | 2. Discussion .....                                 | 47 |
| A. Progress toward on-chip interferometry .....      | 16 | E. Concluding remarks and outlook .....             | 49 |
| B. Precision sensing .....                           | 17 | IX. ATOMTRONICS ENABLED QUANTUM DEVICES             |    |
| C. Cryogenic atom chips and hybrid quantum           |    | AND SENSORS .....                                   | 49 |
| systems .....  | 18 | A. Diodes, transistors, and other discrete          |    |
| D. Concluding remarks and outlook .....              | 20 | components .....                                    | 50 |
| V. QUENCH DYNAMICS OF INTEGRABLE MANY-               |    | B. Atomtronic SQUIDS .....                          | 51 |
| BODY SYSTEMS .....                                   | 20 | C. Sagnac interferometry and rotation sensing ..... | 52 |
| A. Quench dynamics .....                             | 21 | D. Magnetometry .....                               | 53 |
| B. Evolution under integrable Hamiltonians .....     | 21 | E. Concluding remarks and outlook .....             | 54 |
| 1. The Lieb-Liniger model .....                      | 21 | X. TWO LEVEL QUANTUM DYNAMICS IN RING-              |    |
| 2. The XXZ Heisenberg spin chain .....               | 26 | SHAPED CONDENSATES AND MACROSCOPIC                  |    |
| C. Concluding remarks and outlook .....              | 26 | QUANTUM COHERENCE .....                             | 56 |
| VII. NONEQUILIBRIUM PROTOCOLS FOR ONE                |    | A. The Atomtronic quantum interference device:      |    |
| DIMENSIONAL BOSE GASES IN ATOMTRONIC                 |    | AQUID .....   | 56 |
| CIRCUITS .....                                       | 27 | 1. The rf-AQUID qubit .....                         | 57 |
| A. Quench dynamics in the Lieb-Liniger model .....   | 28 | 2. Atomtronic flux-qubit: Ring lattice              |    |
| 1. The quench problem .....                          | 28 | interrupted with three weak links .....             | 58 |
| 2. The quench action .....                           | 29 | B. Demonstration of the one qubit and two qubit     |    |
| 3. Quenches to the attractive regime .....           | 29 | unitary gates .....                                 | 58 |
| B. Floquet Hamiltonian for the periodically tilted   |    | 1. Single qubit gates .....                         | 59 |
| Lieb-Liniger Model .....                             | 29 | 2. Two-qubit coupling and gates .....               | 59 |
| C. Concluding remarks and outlook .....              | 31 | C. Readout of atomtronic qubits .....               | 59 |
| VIII. PERSISTENT CURRENTS AND VORTICES IN            |    | 1. Interferometric detection of the current states  | 59 |
| ATOMTRONIC CIRCUITS .....                            | 32 | D. Experiment realization of the ring-lattice       |    |
| A. Mechanism for producing flow in a racetrack       |    | potential with weak links .....                     | 62 |
| atom-circuit BEC by stirring .....                   | 33 | 1. A ring lattice with single weak link .....       | 62 |
| 1. How stirring a racetrack atom circuit             |    | 2. Experiment realization of the ring-lattice       |    |
| produces flow .....                                  | 33 | potential with three weak links .....               | 62 |
| 2. Creation of a single unit of flow: Vortex         |    | E. Setup for adjustable ring-ring coupling .....    | 63 |
| swap .....   | 33 | F. First experimental demonstration of the          |    |
| 3. Final flow production: Flow overtakes the         |    | interference of atomtronic currents .....           | 64 |
| barrier .....  | 33 | G. Concluding remarks and outlook .....             | 66 |
| B. Persistent currents in coplanar double-ring       |    | XI. TRANSPORT AND DISSIPATION IN ULTRACOLD          |    |
| geometries .....                                     | 34 | FERMI GASES .....                                   | 67 |
| 1. Spontaneous persistent current formation in       |    | A. Superfluid transport with Fermi gases .....      | 68 |
| a ring trap .....                                    | 34 | 1. Fermionic superfluidity and critical velocity .. | 68 |
| 2. Spontaneous persistent current formation          |    | 2. Josephson currents .....                         | 68 |
| in coplanar connected ring traps .....               | 34 | B. Fermionic transport in mesoscopic channels ..... | 69 |
| C. Persistent currents in transversally coupled      |    | C. Fast spin drag in normal Fermi gases .....       | 70 |
| atomtronic circuits .....                            | 36 | D. Concluding remarks and outlook .....             | 71 |
| D. Concluding remarks and outlook .....              | 38 | XII. TRANSPORT IN BOSONIC CIRCUITS .....            | 72 |
| VIII. PHASE SLIP DYNAMICS ACROSS JOSEPHSON           |    | A. Matterwave guides .....                          | 72 |
| JUNCTIONS .....                                      | 39 | B. Ring-leads system .....                          | 75 |
| A. Critical transport and vortex dynamics in a thin  |    | C. Y-junctions .....                                | 75 |
| atomic Josephson junction .....                      | 39 | D. Differences between fermions and hard-core       |    |
| B. Bose-Josephson junction among two one-            |    | boson .....   | 76 |
| dimensional atomic gases: A quantum impurity         |    | E. Entangled state generation with topological      |    |
| problem .....  | 42 | pumping in ring circuits .....                      | 77 |
| 1. Model .....                                       | 42 | F. Concluding remarks and outlook .....             | 77 |
| C. Bose-Einstein condensate confined in a 1D ring    |    | XIII. ARTIFICIAL QUANTUM MATTER IN LADDER           |    |
| stirred with a rotating delta link .....             | 43 | GEOMETRIES .....                                    | 77 |
| 1. Model .....                                       | 44 | A. A boson ring ladder at weak interactions .....   | 78 |

|   |    |
|---|----|
| 1. Model .....  | 78 |
| 2. Ground state of weakly interacting ring .....                                    | 78 |
| 3. Excitation spectrum of weakly interacting ring .....                             | 78 |
| B. The boson ladder at strong interaction .....                                     | 79 |
| C. Ultracold atoms carrying orbital angular momentum (OAM) in a diamond chain ..... | 80 |
| 1. Topological edge states and Aharonov–Bohm caging .....                           | 80 |
| 2. Simulating quantum magnetism with strongly interacting ultracold bosons .....    | 82 |
| D. Concluding remarks and outlook .....   | 83 |
| XIV. QUANTUM-ENHANCED ATOMTRONICS WITH BRIGHT SOLITONS .....                        | 83 |
| A. Scattering properties of attractive bosons against a barrier .....               | 83 |
| B. Creation and manipulation of quantum solitons ..                                 | 84 |
| 1. Quantum solitons in the Bose–Hubbard model .....                                 | 84 |
| 2. Solitons in rotation .....   | 85 |
| 3. Entangling solitons with different $L_z$ .....                                   | 87 |
| C. Concluding remarks and outlook .....   | 87 |
| XV. ATOMTRONICS WITH ALKALINE-EARTH-LIKE METAL ATOMS .....                          | 87 |
| A. Why alkaline-earth-like metal atoms? .....                                       | 87 |
| B. Effective Abelian and non-Abelian gauge fields ..                                | 88 |
| C. Persistent current of $SU(N)$ fermions .....                                     | 89 |
| D. Concluding remarks and outlook .....   | 90 |
| XVI. MANIPULATING RYDBERG ATOMS .....   | 91 |
| A. Driven-dissipative Rydberg systems .....   | 91 |
| B. Microwave-optical conversion using Rydberg atoms .....                           | 92 |
| C. Concluding remarks and outlook .....   | 93 |

## I. INTRODUCTION

Quantum technologies are enabling important innovations in the 21st century with applications in areas as diverse as computation, simulation, sensing, and communication. The core of these new technological developments is the ability to control quantum systems all the way from the macroscopic scale down to the single quantum level. The latter has been achieved in physical systems ranging from atomic and spin systems to artificial atoms in the form of superconducting circuits.<sup>1,2</sup>

In this article, we mostly focus on cold atom systems, where recent technological developments have delivered a collection of magnetic or laser-generated networks and guides in which atomic matter-waves can be controlled and manipulated coherently.<sup>3,4</sup> *Atomtronics* exploits the state of the art in this field to realize matter-wave circuits of ultracold atoms.<sup>4,5</sup> Some key aspects of this emerging field give atomtronic circuits great promise as a quantum technology. First, since atomtronic circuits employ matter-waves of neutral atoms, spurious circuit-environment interactions, which might, e.g., lead to decoherence, are expected to be less serious than in networks employing electrically charged fluids sensitive to Coulomb forces. Second, atomtronic networks can realize new types of circuits with current carriers having bosonic and/or fermionic quantum statistics along with tunable

particle–particle interactions ranging from short-range to long-distance and from attractive to repulsive. Third, recent progress in the manipulation of optical guiding potentials enables engineering of time-dependent circuits whose topology can be reconfigured while they operate.<sup>3,6–9</sup>

The name *Atomtronics* is inspired by the analogy between circuits with ultracold atomic currents and those formed by electron-based networks of conductors, semiconductors, or superconductors. For example, a Bose–Einstein condensate (BEC) confined in a linear optical lattice with a suitable abrupt variation of the particle density can exhibit behavior very similar to that of an electronic diode.<sup>10,11</sup> As another example, a BEC in suitable optical ring trap is the atomic counterpart of the superconducting SQUID of quantum electronics,<sup>11–13</sup> displaying the SQUID’s defining characteristics of quantum interference<sup>15</sup> and hysteresis.<sup>16</sup> It is important to note that because atomtronics is entirely based on flexible potential landscapes and not limited to material properties, it is expected to be possible to create quantum devices and simulators with new architectures and functionalities that have no analog in conventional electronics.

The quantum nature of ultracold atoms as coherent matter waves enables interferometric precision measurements and new platforms for quantum information processing with applications in fundamental science and technology.<sup>17,18</sup> At the same time, atomtronic circuits can serve as powerful probes of many-body quantum regimes: analogous to solid state I–V characteristics, and many-body cold atom systems can be probed by monitoring the current flowing in them while changes are made to external parameters and applied (effective) fields. In this way, atomtronic platforms can be thought of as extensions to the scope of conventional quantum simulators, revisiting textbook scenarios in many-body physics, such as frustration effects, topological constraints, and edge state formation, with the advantages of tunable boundary conditions and minimal finite size effects. Another interesting domain in which atomtronics can play an important role is mesoscopic physics.<sup>19–21</sup> Important themes in the field of mesoscopic physics, such as persistent currents in ring-shaped structures and problems of quantum coherent transport, can be explored with a new twist.

For the implementation of the program sketched above, an important challenge to face in the years to come is to optimize the control of the matter-wave currents in complex networks as, for example, optical lattices, guiding circuits for matter waves based on optical or magnetic fields, or cold atoms–solid state hybrid circuits. On one hand, such a step would be instrumental to harness current and transport for investigations on quantum many-body physics and artificial matter in both the static and dynamic conditions. In particular, Rydberg atoms and ultracold fermionic systems with  $SU(N)$  symmetry provide novel interesting directions to go to. Experimental challenges for this goal are to design improved schemes for controlling the resulting matter-wave interactions and for including advanced schemes for their detection. On the other hand, the control of complex quantum networks would be opening the way to work out new types of devices based on integrated atomtronic circuits. In particular, new chips integrating different technologies, for example, silicon-based electronics and the various atomtronics approaches, would provide a milestone in quantum technology. Concerning potential applications, a certainly important direction pursued in the current research in *Atomtronics* is devoted to interferometry and inertial sensing with enhanced



performance, but quantum simulation and computation as well as all other aspects of quantum technology are accessible. In this context, stabilizing the atomic coherence on small-to-intermediate spatial scales, for example, by smoothing the wave guides, are important challenges to be solved in order to harness the full power of cold-matter-wave quantum technology.

In this review, we summarize recent activities in *Atomtronics* and discuss the future of the field. In Secs. II–IV, we review fabrication principles for atomtronic platforms, ranging from reconfigurable optical potentials employing acousto-optic deflectors, digital micromirror devices, and liquid-crystal spatial light modulators to micro-optical systems and hybrid solid state–cold atom systems circuits where a scanning focused laser beam modifies the current density of a superconducting chip to create the desired trapping potential. These new capabilities open the way to addressing the dynamics of many-body systems, as described in Secs. V and VI. Sections VII and VIII deal with persistent currents in toroidal and ring-shaped condensates. These systems, the simplest atomtronic circuits with a closed architecture, enable the study of basic questions in many-body physics in a variety of new and different conditions. Atomtronic quantum sensors and devices are discussed in Sec. IX. Ring-shaped bosonic circuits are investigated as ideal platforms for matter-wave SQUIDs [the Atomtronic QUantum Interference Device (AQUID)] and flux qubits in Sec. X. These studies have also touched upon a number of fundamental questions, such as macroscopic quantum coherence, the nature of superfluidity in restricted geometries, and vortex dynamics. Transport in fermionic and bosonic circuits are discussed in Sec. XI and XII, respectively. Section XIII deals with bosonic ladders. In addition to their potential relevance to basic research in many-body physics, we envisage that they will be instrumental to the fabrication of coupled atomtronic circuits. In Sec. XIV, we discuss atomtronic circuits that exploit bright solitons both for studying fundamental questions in many-body quantum dynamics and for realizing quantum devices with enhanced performances. Sections XV and XVI deal with alkali-earth atoms with  $SU(N)$  symmetry and Rydberg atoms. To date, the latter have received little attention, but we believe that they offer great promise as an atomtronic quantum technology.

The present article was inspired by the *Atomtronics@Benasque* conference series. The Benasque staff is warmly acknowledged for their invaluable help in the organization of these workshops, and we thank the Benasque director Jose-Ignacio Latorre for his constant support of this line of research.

## II. DYNAMICALLY SCULPTED LIGHT

**M. Baker, G. Gauthier, T. W. Neely, H. Rubinsztein-Dunlop, F. Tosto, R. Dumke, P. Ireland, D. Cassettari**

In recent years, many experiments have been carried out with cold neutral atoms in arbitrary, reconfigurable optical potentials. Single atoms have been trapped in arbitrary-shaped arrays,<sup>22–26</sup> which have subsequently led to the demonstration of topological phases of interacting bosons in one-dimensional lattices.<sup>27</sup> Various configurations of atomtronic circuits have been demonstrated, namely, closed waveguides and Y-junctions,<sup>28</sup> oscillator circuits,<sup>29</sup> atomtronic transistors,<sup>30</sup> rings, and atomtronic SQUIDs (AQUIDs).<sup>13,31,32</sup> Reconfigurable optical potentials have also been used to realize Josephson junctions in

rubidium condensates<sup>12</sup> and in fermionic lithium superfluids in the BCS-BEC crossover.<sup>33</sup> They have even been used for the optimization of rapid cooling to quantum degeneracy.<sup>34</sup> Finally, another area of interest is the realization and study of quantum gases in uniform potentials.<sup>35,36</sup> Some of these experiments are described in detail in Secs. II A–II E.

Static holographic potentials, as opposed to reconfigurable, also play an important role in atomtronics and have been implemented with great success.<sup>37–42</sup> In particular, static holograms can provide substantial advantages for the generation of Laguerre–Gaussian and higher order Hermite–Gaussian modes.<sup>40,43,44</sup> Static hologram techniques, such as optical nanofiber evanescent wave trapping,<sup>45</sup> structured nanosurfaces to create trapping potentials,<sup>46,47</sup> and the use of engineered quantum forces<sup>48</sup> (also known as London–van der Waals or Casimir forces), are promisingly emerging technologies that will benefit the field of atomtronics. However, this section focuses on recently adopted *dynamic* technologies that have opened new avenues of research.

More generally, we note that sculpted light has many more applications beyond cold atom physics, e.g., to microscopy, optical tweezers, and quantum information processing with photonic systems.<sup>3</sup> In this section, we review the tools and techniques that underpin all these experiments: scanning acousto-optic deflectors (AODs), digital micromirror devices (DMDs), and liquid-crystal spatial light modulators (SLMs).

### A. Fast-scanning AODs

By rapidly scanning a trapping laser beam much faster than the trapping frequencies for the atoms, the atoms experience the time-average of the optical potential. Under these conditions, despite the modulated scanning action of the beam, the density of the atom cloud remains constant in time. The spatial location of the beam can be scanned in arbitrary 2D patterns, “painting” the potential landscape, simply by modulating the RF frequencies driving the crystal.<sup>7</sup> The control over the RF power at each scan location allows local control over the potential depth. This feature can be used to error-correct, ensuring smooth homogeneous potentials, or can be deliberately engineered to implement barriers, wells, or gradients in the trap. The trapping geometry can be dynamically changed with the use of deep-memory arbitrary waveform generators or field programmable gate array (FPGA) technology, which combined with nondestructive measurement allows for the real-time correction of the potential. Given the weak axial confinement provided by the scanned beam, this is best used in conjunction with an orthogonal light sheet, which provides tight confinement along the axis of the scanned beam, and ensuring excitation and phase fluctuations in the axial dimension is minimized.<sup>7</sup>

### 1. Feed-forward control

The diffraction efficiency of AODs can change with the drive frequency. In order to correct this, it is generally necessary to use feed-forward to compensate by adjusting the RF power of the AOD crystal and, hence, the beam intensity at each (x,y) location. To correct imperfections in other elements of the trapping potential, one can measure the atomic density distribution in the trap using absorption imaging and apply iterative correction to the RF power at each (x,y) location.<sup>32</sup>

## 2. Phase evolution in time-averaged potentials

A full treatment time-averaged potentials need to include the phase evolution of the condensate under the effect of the scanning beam. The time-varying potential  $V(x, y, t)$  acts to imprint a phase  $\phi$  with the evolution  $\hbar \partial \phi(x, y, t) / \partial t = V(x, y, t)$ . For sufficiently fast scan rates, the imprinted phase effect is negligible, but at slower scan rates, this phase imprinting action can accumulate local phase, leading to residual micromotion in the condensate, and the signatures of which have been observed.<sup>49</sup> This is an important consideration for atomtronic applications where the phase is an observable of interest, such as for guided Sagnac interferometry.<sup>50</sup>

## 3. Atomtronics with time-averaged optical traps

The time-averaged optical dipole traps are extremely versatile, allowing a variety of geometries to be generated, and dynamically changed in structure by real-time adjustment of the scanning pattern. In the context of atomtronic geometries, BECs have been trapped into flat bottom line-traps, rings,<sup>32,51</sup> lattices,<sup>52</sup> and dumbbell reservoirs (Fig. 1). Additionally, single mode matter-wave propagation and coherent phase splitting have been demonstrated in circuit elements such as waveguides and beamsplitters.<sup>28</sup> The time-averaged optical beams can be used to introduce multiple repulsive barriers and stirring elements to study persistent currents and superfluid transport in atomtronic circuits.<sup>12</sup>

## B. Optical potentials with liquid-crystal SLMs

A liquid-crystal SLM spatially modulates the phase of the light. The phase pattern on the SLM acts as a generalized diffraction grating so that in the far field, an intensity pattern is formed, which is used to trap atoms. In practice, the far field is obtained by focusing the light with a lens so that the intensity pattern that traps the atoms is created in a well-defined “output plane” coinciding with the lens focal plane. The SLM acts effectively as a computer-generated hologram, and the light field in the output plane is the Fourier transform of the light field in the SLM plane.

The first experiments with these holographic traps go back over ten years ago.<sup>53,54</sup> A reason for the use of phase-only SLMs, rather than amplitude modulators, is that the former does not remove light from the incident beam. This is advantageous from the point of view of light-utilization efficiency. Moreover, as is shown below, a phase-

only SLM allows the control of both the amplitude and phase on the output plane.

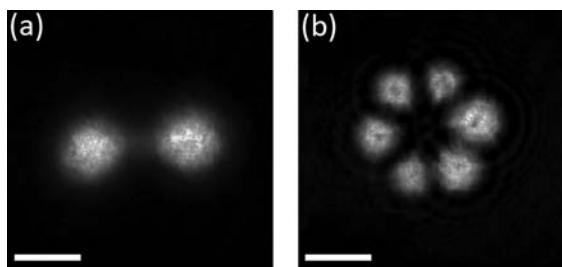
The calculation of the appropriate phase modulation to give the required output field is a well-known inverse problem, which, in general, requires numerical solution. Iterative Fourier Transform Algorithms (IFTAs) are commonly used, and variants that control both phase and amplitude have been recently demonstrated.<sup>55,56</sup> The removal of the singularities (e.g., vortices), which particular pattern optimization techniques can introduce, is widely researched due to their importance for controlled beam shaping<sup>57–60</sup> and, in particular, to confine BECs in uniform potentials.<sup>35,61</sup> One such example is a conjugate gradient minimization technique that efficiently minimizes a specified cost function.<sup>60,62</sup> The cost function can be defined to reflect the requirements of the chosen light pattern, such as removing optical vortices from the region of interest.

The intensity patterns obtained with this method are shown in the first row of Fig. 2. They are taken at a wavelength of 1064 nm, i.e., red-detuned relative to the rubidium transition, causing rubidium atoms to be trapped in the regions of high intensity. The SLM light is focused on the atoms by  $f = 40$  mm lens, giving a diffraction limit of the optical system of  $6 \mu\text{m}$  at 1064 nm.

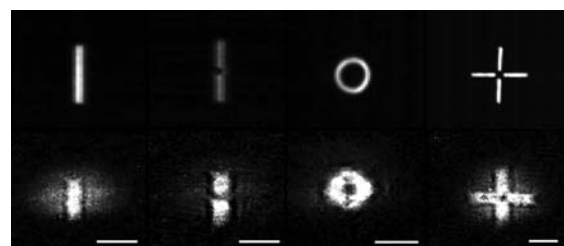
Going from left to right in Fig. 2, shown are a simple waveguide, a waveguide with a potential barrier halfway across, a ring trap, and a crosslike pattern. The latter has been proposed for the study of the topological Kondo effect.<sup>63</sup> In all these light patterns, the phase is constrained by the algorithm. For the simple waveguide, the ring and the cross, a flat phase is programmed across the whole pattern. Controlling the phase this way leads to a well maintained intensity profile shape as it propagates out of the focal plane for up to  $\sim 10$  times the Rayleigh range. By comparison, a pattern with random phase loses its shape much sooner.

Differently from the other three patterns, for the waveguide with the barrier, a sharp  $\pi$  phase change halfway across the line was programmed. In the resulting intensity profile, this phase discontinuity causes the intensity to vanish, hence creating the potential barrier whose width is close to the diffraction limit.

The second row of Fig. 2 shows Rb BECs trapped in the potential created by the SLM light patterns combined with an orthogonal light sheet that provides tight confinement along the axis of propagation of the SLM light.<sup>64,65</sup> The clouds are imaged after a 2 ms time of flight and undergo mean-field expansion during this time, leading to a final density distribution that is more spread out compared to the transverse size of the SLM traps.



**Fig. 1.** Example geometries of time averaged optical dipole potentials: (a) BEC trapped in dumbbell potential with two reservoirs connected through a channel of tunable length and width and (b) ring lattice of BECs. The scale bar on each image indicates  $50 \mu\text{m}$ .



**Fig. 2.**  $^{87}\text{Rb}$  condensates in SLM-generated optical potentials. The top row shows the intensity patterns used for trapping, and the bottom row shows the condensates after a 2 ms time of flight. The ring trap contains  $\sim 10^6$  atoms, while the other traps contain  $\sim 5 \times 10^5$  atoms. The scale bar on the images indicates  $100 \mu\text{m}$ .

Controlling the phase of the light pattern opens new possibilities for the trapping and manipulation of ultracold atoms. Here, we have shown that the phase control gives an alternative way to create barriers close to the diffraction limit by using discrete phase jumps. Liquid-crystal SLMs were also used to transfer phase structure in a four-wave-mixing process in rubidium vapor, in particular trans-spectral orbital angular momentum transfer from near-infrared pump light to blue light.<sup>66</sup> Additionally, they have enabled research into uniform 3D condensates. More recently, they were used in the realization of bottle beams that have been used to create 3D optical trapping potentials for confining Rydberg atoms.<sup>26</sup> In addition to this, the phase control can also be useful for many atomtronics applications, for instance, phase imprint via a Raman transition,<sup>67</sup> and the realization of artificial gauge fields.<sup>68,69</sup>

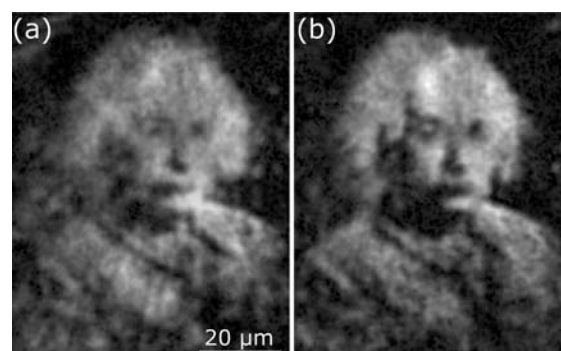
### C. Direct imaged DMD optical potentials

A recent addition to the spatial light modulator family is the digital micromirror device (DMD). Developed for digital light processing (DLP) applications, DMDs consist of millions of individually addressable, highly reflective mirrors. Each hinged mirror, of typical size  $7.56\text{--}10.8\text{ }\mu\text{m}$ , is mounted on a silicon substrate on top of control electrodes. The application of a control voltage tilts the mirrors between two “on” or “off” angles, typically  $\pm 12^\circ$ . The mirror array acts as a dynamical configurable amplitude mask for light reflected from their surface. The DMDs can be placed in the Fourier plane of the imaging/project system, similar to typical phase-based SLMs, where it can modulate both the phase and the amplitude of the light.<sup>72</sup> If phase modulation is not required, the DMD can be used as a binary amplitude mask in the object plane, similar to its DLP applications.<sup>9,73</sup> In the “DC” mode, the mirrors are fixed to the on angle and a static pattern can be projected. The true versatility of the device, however, lies in its dynamical (“AC”) capability with full frame refresh rates exceeding 20 kHz.

#### 1. Half-toning and time-averaging

The projected image from the DMD is binary in nature. Although this would appear as a significant limitation in producing arbitrary optical potentials, a number of techniques exist to overcome this issue. The first of these is half-toning, or error-diffusion, which takes advantage of the finite optical resolution of the projection optical system to increase the amplitude control. With suitable high magnification, such that the projected mirror size is smaller than the resolution, multiple mirrors contribute to each resolution spot in the projected plane.<sup>74</sup> In this way, half-toning can be used to create intensity gradients in the light field, as shown in Fig. 3(a). Same as in the case of time-averaged AOD traps, feed-forward using the atomic density<sup>32,70,71</sup> can be performed to correct for imperfections in the projection potential, as shown in Fig. 3(b).<sup>71</sup>

One can also make use of the high-speed modulation of the mirrors to further improve the intensity control. The mirror array of the DMD is capable of switching speeds from DC to 20 kHz. By varying the on/off time of individual mirrors (pulse-width modulation), the time-average of the resulting light field can be utilized to improve the smoothness of the projected potentials.<sup>71</sup>



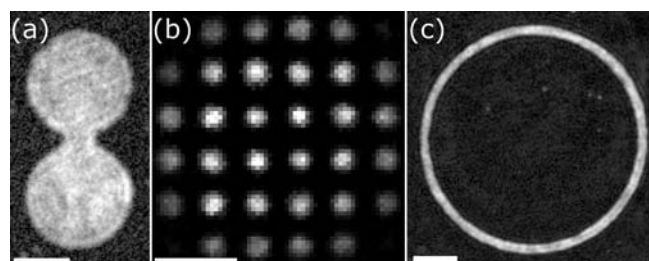
**Fig. 3.** Generating complex potentials using half-toning. (a) Initial *in situ* image of the BEC density in a half-toned potential of Einstein calculated using the optical system parameters. (b) Final converged BEC image after 11 feedback iterations where the atomic density is used to iteratively correct for imperfection in the density generated using the method described by Tajik *et al.* (Ref. 70) and Gauthier (Ref. 71).

### 2. Atomtronics with DMDs

Atomtronics studies how to use neutral atom currents to create circuits that have properties similar to existing electrical devices. The advances in control and increased resolution of trapping potentials have been instrumental in the development of this field. The dynamic control over the potential given by DMDs have allowed time dependent implementations. Combined with other techniques such as the optical accordion lattice,<sup>73</sup> which allows smooth transitions between quasi-2D and 3D systems, they open up further avenues of control for future studies. The high resolution projection of DMD optical potentials enables the creation of complex masks. These have facilitated the study of superfluid transport in a variety of traps. Figure 4 shows three relevant geometries for superfluid transport studies.

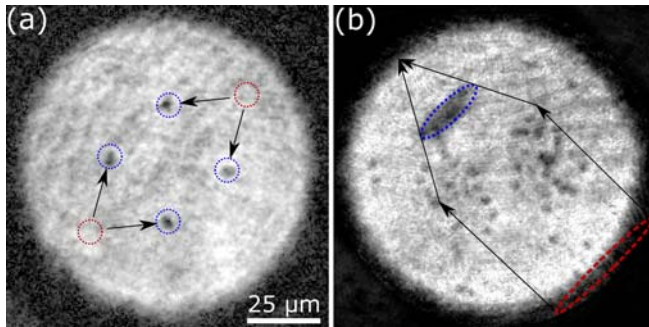
### 3. Turbulence with DMDs

The dynamic properties of DMDs can be used for the creation of turbulence. As shown in Fig. 5, these techniques have been used to study Onsager vortices and their emergence in superfluids,<sup>75,76</sup> the



**Fig. 4.** Useful atomtronics geometries created with directly imaged DMD trapping. (a) A dumbbell geometry of two reservoirs connected by a channel where the reservoir size, channel length, and channel width can be varied to study superfluid transport. (b) Square lattice of BECs with a period of  $10\text{ }\mu\text{m}$  formed using the projection of a half-toned DMD pattern. The lattice period can be dynamically increased or decreased. (c) A ring-shaped BEC with a diameter of  $110\text{ }\mu\text{m}$  and a radial width of  $10\text{ }\mu\text{m}$  is useful for interferometry and transport experiments. The scale bar on each image indicates  $20\text{ }\mu\text{m}$ .





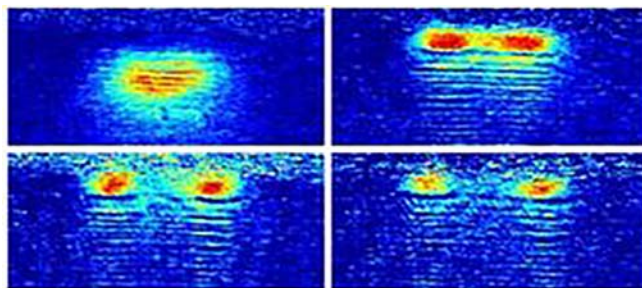
**FIG. 5.** Creating vortices using the dynamical capabilities of DMDs. (a) Deterministic creation of vortices using the “chopstick” method outlined in Samson *et al.*<sup>79</sup> The dashed red (blue) circle represent the initial (final) position of the optical barriers with arrows indicating their trajectories. After a short 3 ms time of flight, the vortices are visible as density dips (black dots). (b) Creation of a vortex cluster, similar to the procedure used by Gauthier *et al.* (Ref. 75) and Johnstone *et al.* (Ref. 76), by sweeping a paddle-shaped optical barrier through a condensate. The red (blue) dashed ovals represent the paddle position at initial (picture) time with the arrows indicating the trajectory of the edges of the paddle.

creation of tunable velocity solitons,<sup>77</sup> and equilibration of chiral vortex clusters.<sup>78</sup>

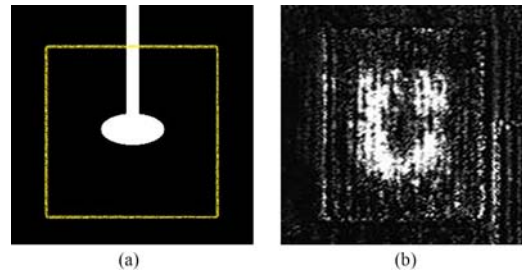
#### D. Hybrid atomic-superconducting quantum systems

Superconducting (SC) atom chips have significant advantages in realizing trapping structures for ultracold atoms compared to conventional atom chips.<sup>4,80–85</sup> These advantages have been extended further by the development of the ability to dynamically tailor the superconducting trap architecture. This is done by modifying the current density distribution in the SC film through local heating of the film using dynamically shaped optical fields. This allows for the creation of desired magnetic trapping potentials without having to change the chip or the applied electrical field.

Typically, a high-power laser and a DMD are used to create and shape the light field used to destroy the superconductivity and influence the shape and structure of a trap. Various trapping potentials have been realized using this technique, in particular, to split a single trap (see Fig. 6) or to transform it into a crescent or a ringlike trap (see Fig. 7). Since the atomic cloud evolves with the trapping potential, cold atoms can be used as a sensitive probe to examine the real-time



**FIG. 6.** Absorption images of the atomic cloud splitting taken after 0 (top-left), 20 (top-right), 30 (bottom-left), and 40 (bottom-right) ms of illumination time, respectively.



**FIG. 7.** DMD image on the left, where the real dimension of the SC film is highlighted in yellow and absorption image of the atomic cloud on the right.

magnetic field and vortex distribution. Simulations of the film heating, the corresponding redistribution of sheet current density, and the induced trapping potentials have been found to agree closely with experiments. Such simulations help us to better understand the process and can be used to design traps with the needed properties.

More complex structures can be achieved by increasing the heating pattern resolution. This method can be used to create magnetic trap lattices for ultracold atoms in quantum computing applications and, in particular, optically manipulated SC chips open new possibilities for ultracold atoms trapping and design of compact on-chip devices for investigation of quantum processes and applications in atomtronics.<sup>86–92</sup>

#### E. Concluding remarks and outlook

In this section, we have described the suite of technologies available to the experimenter for creating configurable optical potentials for ultracold atoms, primarily discussing AOMs, SLMs, and DMDs. A hybrid technique utilizing optical configurable potentials to shape magnetic potentials through superconducting quantum chips intermediary were also discussed. These technologies have drastically improved the control and manipulation of ultracold neutral atoms.

Although previously available static holograms technologies provided great control for the creation of optical potentials and are still usually better for 3D trapping potentials, the dynamic manipulation capabilities presented here have enabled new classes of experiments with ultracold atoms. For example, dynamically modulated DMDs have facilitated new studies of two-dimensional-quantum turbulence<sup>75,76,78</sup> and condensate evolution in response to rapidly quenched trapping potentials.<sup>31,96</sup> AOMs have enabled steerable arrays of single atoms,<sup>23,52,94</sup> facilitating quantum simulation experiments. Furthermore, the rapid reconfigurability of DMD traps has enabled groundbreaking studies in the emerging field of atomtronics, where the system parameters can be easily tuned.<sup>29,33,77,95,99</sup>

As the technology behind optical manipulation continues to mature and evolve through the increase in SLMs pixel array sizes and switching frequencies, these sculpted light and hybrid techniques are sure to have an even bigger impact on the development of atomtronics.

#### ACKNOWLEDGMENTS

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### III. IMPLEMENTING RING CONDENSATES

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The many interesting properties of degenerate quantum gases, such as phase coherence, superfluidity, and vortices, naturally make the geometry of these systems of great interest. Ring systems are of particular interest, as the simplest multiply connected geometry for coherent matter-wave guiding and as a potential building block for circuitual atomtronic devices. In addition, ring systems have interesting properties such as persistent flow, quantum hall states, and the potential for Sagnac interferometry.

Advances in the control of quantum gases have seen the development of atom waveguides formed from both magnetic trapping and magnetic resonance, and optical dipole trapping, and more recent implementations using hybrids of both. These approaches satisfy the criteria needed for coherent quantum matter-wave flow: i.e., the waveguides are *smooth* and can form *loops* that are *dynamically* controllable.

#### A. General features of ring traps

Irrespective of the mechanism of trapping, magnetic or optical, some common parameters for ring traps can be described. We will restrict our discussion to ring traps that can be considered approximately harmonic; in cylindrical co-ordinates, the ring potential with radius  $R$  is expressed in terms of radial and vertical trapping frequencies  $\omega_\rho$  and  $\omega_z$ , respectively,

$$V(\rho, z) = \frac{1}{2} m \omega_\rho^2 (\rho - R)^2 + \frac{1}{2} m \omega_z^2 z^2. \quad (1)$$

Considering now a trapped gas within this potential, the connected geometry of the ring trap results in modifications to the usual derivation for the condensate critical temperature  $T_c$  for a 3D harmonically trapped gas, yielding<sup>97</sup>

$$T_c = \left( \frac{\sqrt{2} N_0 \hbar^3 \omega_\rho \omega_z}{1.514 k_b^{5/2} m^{1/2} \pi R} \right)^{2/5}, \quad (2)$$

where  $N_0$  is the atom number. For sufficiently elongated geometries, such as cigar traps, or ring traps with long azimuthal length, a regime of thermally driven phase-fluctuations in the condensate can exist<sup>98,102</sup> even at temperatures below  $T_c$ . These phase-fluctuations are suppressed when the correlation length is larger than the system size, which for a ring geometry is half the azimuthal circumference, or  $\pi R$ . As we are typically interested in fully phase coherent ring traps, we can define this transition temperature  $T_\phi$ ,<sup>97</sup>

$$T_\phi = \frac{\hbar^2 N_0}{k_b m \pi R^2}. \quad (3)$$

Finally, in the Thomas–Fermi approximation, where the interaction energy dominates, the chemical potential in the ring trap can be expressed in terms of the trapping parameters,<sup>100</sup>

$$\mu = \hbar \sqrt{\frac{2 N_0 \omega_\rho \omega_z a_s}{\pi R}}, \quad (4)$$

and  $a_s$  is the s-wave scattering length.

In this section, we will discuss the experimental and theoretical developments in all three types of waveguide approach. In what follows, in Sec. III B, we discuss approaches primarily involving magnetic and radio-frequency fields; and in Sec. III C, we will discuss optical and hybrid approaches to implementing ultracold atoms and condensates in rings before concluding in Sec. III E.

#### B. Techniques based on magnetic traps

Experimental techniques for trapping atoms in magnetic fields are well developed since the first BECs, and it is natural to consider such an approach, and build on that approach, to make ring waveguides. Nevertheless, this brings particular challenges because of the need to satisfy Maxwell's equations for fields trapping in a ring geometry, the need to avoid the loss of atoms from Majorana spin flips, occurring in the vicinity of field zeros, and the desire, for some experiments, to have trapping systems with high symmetry.

The earliest examples of waveguides for ultracold atoms were produced using static magnetic fields, where DC current carrying wires were used to create large area ring<sup>101</sup> and stadium<sup>102</sup> geometries which initially trapped thermal atoms. With Ref. 103, we had the first demonstrations of a ring waveguide with a Bose-condensed gas. Subsequent experimental developments can be divided into systems which principally use macroscopic coils for generating the magnetic trap, and those systems which employ microfabricated structures in an *atom chip* to generate the spatially varying potentials. We will briefly discuss the latter next and the former in Secs. III B 1–III B 3.

The appeal of atom-chip traps is their compact footprint, potential portability, and the ability to fabricate quite complex geometries, switches, and antenna components into a compact package.<sup>104,105</sup> Additionally, the close proximity of the wires allows high trapping frequencies to be achieved, even for modest currents. However, trapping in close proximity to a surface brings with it its own challenges. Foremost of these are the corrugations in the magnetic guiding potential, which arise from imperfectly directed currents in the conducting material. An additional challenge is the perturbing effect of the end connections to supply current in and out of the conducting ring. Although these problems can be alleviated to some degree by the use of AC fields,<sup>106</sup> which provides a smooth time-averaged current in the wire, as well as switching elements at the end connections to minimize the perturbative phase effects on the ring condensate,<sup>107</sup> they cannot be removed completely. A comprehensive survey on the implementation of ring traps based on atom-chips, and their applications, is covered in detail in Ref. 108.

Here, we will focus our attention on ring traps derived from a combination of static magnetic traps with RF and modulated fields. Using macroscopically large conducting elements requires the use of high currents and occupies a greater size, but there are significant gains in the resulting trap smoothness, as the conducting elements are far from the trapping region. This makes such magnetic traps ideal for producing corrugation free toroidal waveguides for coherent matter, which is detailed in this section.



However, the complexity of the fields requires an atom-chip approach to a pure magnetic waveguide system<sup>104,105</sup> and this brings a difficult problem for the perfect ring waveguide because of the need to get the currents into, and out of, the wires that define the waveguide. We can try to live with this,<sup>108</sup> but asymmetry seems inevitable. We can think of tricks, for example, as the atoms go around the ring, and we can switch the current between different sets of 130 wires as in Ref. 107. This would avoid the bumps and humps in the waveguide, which occur in the places where current enters and leaves the defining structures at the expense of potential losses and heating as the guides are switched over.

### 1. RF dressing and bubbles

It is not obvious that micrometer-scale trapping structures for ultracold atoms can be created using macroscopic scale magnetic coils. However, by means of the addition of radio-frequency coils, magnetic traps with a simple trapping geometry can be transformed into ring traps and other topologies. The theoretical basis is to treat the atom and radio-frequency field with adiabatic following and the dressed-atom theory.<sup>109</sup> Originally introduced in the optical domain by Cohen-Tannoudji and Reynaud,<sup>109</sup> we adapt it here in the radio-frequency domain where it has found several applications (see also Secs. III B 2, III B 3, and III D). The approach is suitable for ultracold atoms in magnetic traps where the trap potential is governed by the spatially varying Zeeman energy and the spatially varying energy difference between Zeeman levels can be in the radio-frequency range.<sup>110,111</sup> The method relies on the adiabatic following of local eigenstates, and it is notable that the superpositions of Zeeman states can provide some resilience to temporal noise and surface roughness.<sup>106</sup> The combination of static magnetic fields and radio-frequency fields with their different spatial and vector variation allows flexibility in the resulting potentials for the creation of shell potentials, rings, tubes, and toroidal surfaces among others.<sup>110,111</sup>

As a simple example, we can consider a simple spatially varying static field and a uniform radio-frequency field. A simple spatially varying magnetic field (obeying Maxwell's equations) is the quadrupole field,

$$\mathbf{B}_0(\mathbf{r}) = b'(x\hat{\mathbf{e}}_x + y\hat{\mathbf{e}}_y - 2z\hat{\mathbf{e}}_z), \quad (5)$$

where  $b'$  represents the gradient of the field in the  $x$ - $y$  plane. This field is often generated by a pair of coils with current circulating in opposite directions. When an atom interacts with this static field via its magnetic dipole moment  $\mu$ , we obtain the ubiquitous interaction energy,

$$U(\mathbf{r}) = -\mu \cdot \mathbf{B}_0(\mathbf{r}) \rightarrow m_F g_F \mu_B |\mathbf{B}_0(\mathbf{r})|, \quad (6)$$

responsible for magnetic potentials and the Zeeman energy splitting. The second form for  $U(\mathbf{r})$  has the integer or half integer  $m_F = -F, \dots, F$ , which arise from the quantization of the energy along with the Landé  $g$ -factor  $g_F$  and Bohr magneton  $\mu_B$ . For our example static field [Eq. (5)], the resulting potential is  $U(\mathbf{r}) = m_F \hbar \alpha \sqrt{x^2 + y^2 + 4z^2}$ , where  $\alpha = g_F \mu_B b' / \hbar$ .

In the next step toward radio-frequency dressed potentials, we add the RF field. The interaction is still given by Eq. (6) but with the replacement  $\mathbf{B}_0(\mathbf{r}) \rightarrow \mathbf{B}_0(\mathbf{r}) + \mathbf{B}_{\text{rf}}(\mathbf{r}, t)$ . The oscillating radio-frequency field  $\mathbf{B}_{\text{rf}}(\mathbf{r}, t)$  is, in general, off-resonant to the local Larmor

frequency or local Zeeman energy spacing  $|g_F \mu_B \mathbf{B}_0(\mathbf{r})|$ , and we define a spatially varying detuning of the RF field as

$$\delta(\mathbf{r}) = \omega_{\text{rf}} - \omega_L(\mathbf{r}). \quad (7)$$

Those locations defined by  $\delta(\mathbf{r}) \rightarrow 0$  typically define a surface in space where RF resonance is found, and correspondingly, there is a minimum in the interaction energy overall.<sup>110,111</sup> In the linear Zeeman regime, the local Larmor frequency is given by

$$\omega_L(\mathbf{r}) = \frac{|g_F \mu_B \mathbf{B}_0(\mathbf{r})|}{\hbar}, \quad (8)$$

which is derived from the static potential  $U(\mathbf{r})$ . The oscillating field  $\mathbf{B}_{\text{rf}}(\mathbf{r}, t)$  yields an interaction energy<sup>110,111</sup> in terms of a Rabi frequency  $\Omega_0(\mathbf{r})$ ,

$$\hbar \Omega_0(\mathbf{r}) = \frac{g_F \mu_B}{2} |\mathbf{B}_{\text{rf}}^\perp(\mathbf{r})|, \quad (9)$$

where the factor of two arises from the rotating wave approximation in the case of linear polarization (more general polarizations are discussed in Ref. 111), and  $\mathbf{B}_{\text{rf}}^\perp(\mathbf{r})$  is the component of  $\mathbf{B}_{\text{rf}}(\mathbf{r}, t)$  perpendicular to the local static field  $\mathbf{B}_0(\mathbf{r})$ . Finally, by combining the energies (6) and (9) through diagonalization of the Hamiltonian in a full treatment,<sup>110,111</sup> we obtain the local eigenenergies, or dressed potentials,

$$U(\mathbf{r}) = m'_F \hbar \sqrt{\delta^2(\mathbf{r}) + \Omega_0^2(\mathbf{r})} \\ = m'_F \sqrt{[\hbar \omega_{\text{rf}} - \hbar \omega_L(\mathbf{r})]^2 + [g_F \mu_B |\mathbf{B}_{\text{rf}}^\perp(\mathbf{r})|/2]^2}, \quad (10)$$

where  $m'_F$  are a set of integers, or half-integers, similar to the  $m_F$  described above.

The result of this is that slow atoms are confined by the potential (10), which in a typical configuration, and to a first approximation, confines atoms to an iso- $B$  surface defined by  $\hbar \omega_{\text{rf}} - \hbar \omega_L(\mathbf{r}) = 0$ , which approximately reduces the value of  $U(\mathbf{r})$  in Eq. (10). The term  $g_F \mu_B |\mathbf{B}_{\text{rf}}^\perp(\mathbf{r})|/2$  also plays a role, and in particular, it can be zero at certain locations on the trapping surface allowing the escape of atoms. This latter effect prevents the trapping of atoms in a shell potential by using the static quadrupole field (5). However, shell potentials are possible with different field arrangements such as those arising from the Ioffe-Pritchard trap and variations,<sup>110-115</sup> which have become candidates for experiments on the International Space Station.<sup>116</sup> The requirement is simple for a local extremum in the *magnitude* of the field  $\mathbf{B}_0(\mathbf{r})$  together with a nonzero  $\mathbf{B}_{\text{rf}}^\perp(\mathbf{r})$ . The reason for the interest in shell potentials in the earth orbit is that on the earth's surface a gravitational term  $mgz$  should be added to Eq. (10), which plays an important role for larger and interesting shells (e.g., see Sec. III B 3).

Although the matter-wave *bubbles* produced by shell potentials have become an object of great interest, the shell potentials themselves are the building blocks for other potentials of interest such as ring traps: we will see an example in Sec. III D. Another example is in Sec. III B 2, where a modulated bias field is used to make a ring trap: then  $\mathbf{B}_0(\mathbf{r}) \rightarrow \mathbf{B}_0(\mathbf{r}) + \mathbf{B}_m(\mathbf{r}, t)$  and  $\mathbf{B}_m(\mathbf{r}, t)$  is a field varying in space, and time, but typically at a frequency rather lower than the radio-frequency case.

### 2. Waveguides formed from time-averaged adiabatic potential (TAAP)

Time averaged adiabatic potentials (TAAPs) allow the generation of extremely smooth matterwave guides,<sup>117</sup> which can be shaped into

a half-moon or ring (see Fig. 8). They are an excellent candidate for matterwave optics, long-distance transport experiments, and interferometry in an atomtronic circuit.<sup>117–119</sup> TAAPs are formed by applying an oscillating homogeneous potential to the adiabatic bubble traps described in Sec. III B 1. If the modulation frequency ( $\omega_m = 2\pi f_m$ ) is small compared to the Larmor frequency, but fast compared to the trapping frequency of the bubble trap, then the effective potential for the atoms is the bubble potential time-averaged over one oscillation period.<sup>120</sup> Let us consider TAAP potentials formed from a quadrupole bubble trap and an oscillating homogeneous field of the form  $\mathbf{B}_m = \{0, 0, B_m \sin \omega_m t\}$ . The modulation field simply displaces the quadrupole (and thus the bubble trap) by  $z_m = \alpha^{-1} B_m \sin \omega_m t$  at an instant in time. In order to find the *effective* potential that the atoms are subjected to by this method, one calculates the time-average. Time-averaging of a concave potential increases the energy of the bottom of the trap, as is readily illustrated by taking the time average of a harmonic potential jumping between two positions: the curvature does not change since it is everywhere the same; however, the energy of the trap bottom increases since it is at exactly the crossing point between the two harmonic potentials. Returning to the modulated bubble trap, one notices that the modulation is orthogonal to the shell at the poles of the shell ( $x = y = 0$ ), but tangential to the shell on the equator ( $z = 0$ ). Therefore, the time averaging causes a larger increase in the trapping potential at the poles rather than the equator and, therefore, creates a ringlike structure.

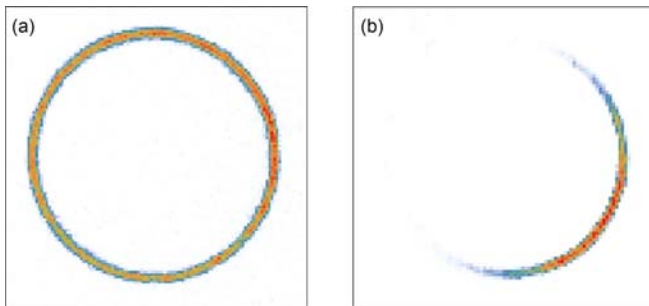
Assuming that  $\omega_{RF}$  is modulated such as to stay resonant on the ring and to keep  $\Omega_{RF}$  constant, the vertical and radial trapping frequencies can be controlled via the relative amplitude of the modulation  $\beta = g_F \mu_B B_m / \hbar \omega_{RF}$  as  $\omega_\rho = \omega_0 (1 + \beta^2)^{-1/4}$  and  $\omega_z = 2\omega_0 \sqrt{1 - (1 + \beta^2)^{-1/2}}$ , where the radial trapping frequency of the bare bubble trap is  $\omega_0 = m_F g_F \mu_B \alpha (m \hbar \Omega_{RF})^{-1/2}$  with the mass of the atom  $m$ , the  $g_F$  is the Landé  $g$ -factor of the considered hyperfine manifold,  $\mu_B$  is the Bohr magneton, and  $\Omega_{RF}$  the Rabi frequency of the dressing RF. In order to achieve large RF field strengths ( $\approx 0.3$ – $1$  G) and Rabi frequencies, ( $\Omega_{RF}$ ), one usually has to use RF-resonators, which make it very difficult to tune the RF frequency, and which results in a somewhat weaker confinement in the axial (i.e., vertical) direction. Trapping frequencies of the order of a hundred hertz are readily achieved.

In many cases, it is also desirable to confine the atoms azimuthally. This is readily achieved either by tilting the ring away from being

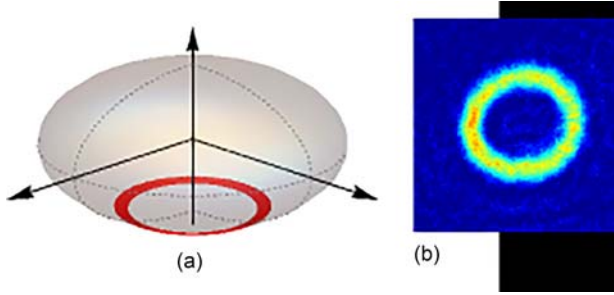
perfectly horizontal or by modifying the polarization of the rf-field. The half-moon shaped BEC in Fig. 8(b) was formed this way. A gravito-magnetic trap results from tilting the direction of the  $B_m$  and thus tilting the ring against gravity.<sup>118</sup> The gravito-magnetic potential forms a single minimum much like a tilted rigid pendulum. One can also create a trap by changing the polarization of the dressing RF: tilting a linear polarization from the  $z$ -axis will cause due to its projection on the local B-field, and a sinusoidal modulation of the Rabi frequency along the ring resulting in a two minima on opposite sides of the ring. Alternatively an elliptical RF polarization creates a single minimum. Combining these modulation techniques permits the creation of two arbitrary placed traps along the ring or, more generally, any longitudinal confinement of the form  $a_1 \sin(\phi + \phi_1) + a_2 \sin(2\phi + \phi_2)$ , where  $\phi$  is the azimuthal angle and  $\phi_1$  and  $\phi_2$  are phase offsets. Note that there are no angular spatial Fourier components higher than  $2\phi$  present in the system.

Thermal atoms and BECs are readily loaded into the gravito-magnetic TAAPs from a trapping-frequency-matched dipole trap. This can be done fully adiabatically by ramping down the dipole confinement and at the same time ramping up the TAAP trap. With a sufficiently high level of control on the rf-fields, one can also load them from a TOP trap via a tilted dumbbell-shaped trap.<sup>118</sup> Once in the ring, one can then manipulate the atoms with a simple manipulation of the time-averaging fields: The depth of the azimuthal trap can be changed by modifying the degree of tilt applied to the modulation field ( $\mathbf{B}_m$ ). By changing the direction of the tilt (i.e., the phase between the modulation fields in the  $x$  and  $y$  directions), one can move the trap along the ring. This can be used, e.g., to accelerate the atoms along the ring with angular momenta of  $40\,000 \hbar$  per atom being readily achieved.<sup>117</sup> They can then travel in the waveguides over distances of tens of centimeters without any additional heating associated with the propagation. One can also remove the azimuthal confinement and allow the condensates to expand around the ring. Viewed in the corotating frame at high angular momenta, the atoms see an exceptionally flat potential with the largest resulting density fluctuations corresponding to an energy difference of a few hundred picokelvin: this is equivalent to a few nanometers in height.<sup>117</sup> Current experiments have been performed with BECs in the Thomas–Fermi regime with about 20 transverse vibration modes occupied. The 1D regime is readily accessible simply by reducing the atom number and increasing the radius of the ring.

The complete lack of any roughness combined with a picokelvin level control of the trapping parameters make the TAAP waveguides a very good candidate for guided matterwave interferometry and the study of ultralow energy phenomena such as long-distance quantum tunneling. A remaining challenge is to completely fill the ring with a phase coherent condensate. Current experiments allowed a condensate to expand along the ring, which converts the chemical potential of the BEC into kinetic energy. When the condensate touches itself at the opposite side of the ring, the two ends have a finite velocity in opposite directions, resulting in a spiral BEC, i.e., a BEC wrapped around itself. Using atom-optical manipulation of the expansion process, kinetic energies in the pico-kelvin range (a few hundred micrometers per second) can readily be achieved. It will be interesting to study the very low energy collisions that will lead to a thermalization of this system. A promising approach for a fully phase-coherent ring-shaped condensate is to first fill a small ring and then increase its radius. This should



**FIG. 8.** (a) BEC in a symmetric, ring-shaped TAAP ring with a diameter of 470  $\mu\text{m}$ . (b) BEC in a circular TAAP matterwave guide with superimposed gravito-magnetic modulation in the azimuthal direction.



**Fig. 9.** (a) Density contour (red annulus) for a BEC rotating at  $1.06 \omega_r$  in the shell trap (gray ellipsoid). (b) *In situ* integrated 2D density of a dynamical ring. Picture size:  $130 \times 130 \mu\text{m}^2$ . Reprinted with permission from Guo *et al.*, Phys. Rev. Lett. **124**, 025301 (2020). Copyright 2020, American Physical Society.

not induce any additional phase fluctuations, despite the fact that the lowest excitation has an energy of  $E = \hbar^2 / (2mr^2)$ , which for a ring of 1 mm radius is 3 fK.

### 3. Dynamical ring in an rf-dressed adiabatic bubble potential

There is a formal analogy between the Hamiltonian of a neutral gas in rotation and the one of a quantum system of charged particles in a magnetic field. This makes rotating superfluids natural candidates to simulate condensed matter problems such as type II superconductors or the quantum Hall effect.<sup>121,122</sup> For a quantum gas confined in a harmonic trap of radial frequency  $\omega_r$  and rotating at angular frequency  $\Omega$  approaching  $\omega_r$ , the ground state of the system reaches the atomic analog of the lowest Landau level (LLL) relevant in the quantum Hall regime.<sup>123–125</sup> Reaching these fast rotation rates is experimentally challenging in a harmonic trap because the radial effective trapping potential in the rotating frame vanishes due to the centrifugal force. To circumvent this limit, higher-order confining potentials have been developed,<sup>126</sup> which allow to access the regime where  $\Omega$  even exceeds  $\omega_r$ .

The adiabatic bubble trap has many features that make it a very good candidate to explore this regime. Indeed, it is very smooth, and easy control of its anisotropy is possible through the dressing field polarization.<sup>127</sup> This allows us to deform the bubble and rotate the deformation around the vertical axis in a very controlled way, allowing us to inject angular momentum into the cloud. The curved geometry of the bubble provides naturally the anharmonicity required to rotate the atoms faster than the trapping frequency  $\omega_r$  at the bottom.

In the experiment at LPL,<sup>128</sup> the atoms are placed in a quadrupole magnetic field of symmetry axis  $z$  dressed by a radio-frequency (rf) field of maximum coupling  $\Omega_0$  at the bottom of the shell. Here, the equilibrium properties in the absence of rotation ( $\Omega = 0$ ) are well known:<sup>127</sup> the minimum of the trapping potential is located at  $r = 0$  and  $z = z_0$ , and around this equilibrium position, the potential is locally harmonic with vertical and radial frequencies  $\omega_z = 2\pi \times 356$  Hz and  $\omega_r = 2\pi \times 34$  Hz without measurable in-plane anisotropy. This trap is loaded with a pure BEC of  $2.5 \times 10^5$   $^{87}\text{Rb}$  atoms with no discernible thermal fraction. This atomic cloud has a chemical potential of  $\mu/\hbar = 2\pi \times 1.8$  kHz, which is much greater than  $\omega_r$  and  $\omega_z$  and well in the three-dimensional Thomas-Fermi (TF) regime. In addition to the dressing field, a radio-frequency knife with

frequency  $\omega_{\text{kn}}$  is used to set the trap depth to approximately  $\omega_{\text{kn}} - \Omega_0$  by outcoupling the most energetic atoms in the direction transverse to the ellipsoid.<sup>129,130</sup>

In a frame rotating at frequency  $\Omega$ , the effective dressed trap potential is the usual trap described above with the addition of a  $-\frac{1}{2}M\Omega^2 r^2$  term taking into account the centrifugal potential. In this frame, the atomic ground state consists of an array of vortices of quantized circulation, each vortex accounting for  $\hbar$  of angular momentum per atom. When only a few vortices are present, the velocity field differs strongly from the one of a classical fluid, but for a sufficiently large number of vortices, the superfluid rotates as a solid body with a rotation rate  $\Omega$ . When  $\Omega < \omega_r$ , the equilibrium position remains on the axis  $r = 0$  at  $z = z_0$ , and the only difference is a renormalization of the radial trapping frequency:  $\omega_r^{\text{eff}} = \sqrt{\omega_r^2 - \Omega^2}$ . Of course, as this frequency decreases, the trap anharmonicity becomes more important in the determination of the cloud shape.

For  $\Omega > \omega_r$ , the trap minimum is located at a nonzero radius. In this situation, a hole grows at the trap center above a critical rotation frequency  $\Omega_h$ ,<sup>131</sup> leading to an annular two-dimensional density profile [Fig. 9(a)], which we will refer to as a “dynamical ring.”<sup>128</sup> Moreover, the velocity of the atomic flow is expected to be supersonic,<sup>132</sup> i.e., exceeding by far the speed of sound. For increasing  $\Omega$ , one expects the annular gas to sustain vortices in its bulk up to a point where the annulus width is too small to host them. The gas should then enter the so-called “giant vortex” regime<sup>132,133</sup> where all the vorticity gathers close to the center of the annulus.

The experimental sequence is the following: angular momentum is injected into the cloud by rotating the trap with an ellipsoidal anisotropy at a frequency  $\Omega = 31$  Hz. The trap rotation is then stopped and isotropy is restored. At this moment, which we take as  $t = 0$ , the cloud shape goes back to circular with an increased radius due to its higher angular momentum. An additional evaporation process, selective in angular momentum, continuously accelerates the superfluid and increases its radius.<sup>128</sup> Due to this size increase, the chemical potential is reduced and the gas enters the quasi-2D regime  $\mu \leq \hbar\omega_z$ . After a few seconds, a density depletion is established at the center of the cloud which is a signature of  $\Omega$  now exceeding  $\omega_r$ . After a boost in selective evaporation due to a lowering of the frequency of the rf knife, a macroscopic hole appears in the profile, indicating that  $\Omega$  is now above  $\Omega_h$  and that a fast rotating dynamical ring with a typical radius of  $\sim 30 \mu\text{m}$  has formed as can be seen in Fig. 9(b). The rotation keeps increasing, and a ring is still observable after  $t = 80$  s. Rotational invariance is critical in that regard and is ensured at the  $10^{-3}$  level by a fine tuning of the dressing field polarization and of the static magnetic field gradients.<sup>111</sup>

A Thomas-Fermi profile convoluted with the imaging resolution is much better at reproducing the experimental density profile than a semiclassical Hartree-Fock profile, demonstrating that the samples are well below the degeneracy temperature. Using the Thomas-Fermi model, we can estimate the properties of the cloud. For example, the ring obtained at  $t = 35$  s has a chemical potential of  $\mu/\hbar \simeq 2\pi \times 84$  Hz and an averaged angular momentum per particle  $\langle \hat{L}_z \rangle / N \simeq \hbar \times 317$ . Interestingly, the estimated peak speed of sound  $c = \sqrt{\mu/M} \simeq 0.62$  mm/s at the peak radius  $r_{\text{peak}}$  is much smaller than the local fluid velocity  $v = \Omega r_{\text{peak}} \simeq 6.9$  mm/s: the superfluid is, therefore, rotating at a supersonic velocity corresponding to a Mach



number of 11. Moreover, due to the continuous acceleration of the rotation, the dynamical ring radius grows gradually with time which results in a decrease in the chemical potential and an increase in the Mach number. For  $t > 45$  s, the chemical potential is below  $2\hbar\omega_r$  and the highest measured Mach number is above 18.

Superfluidity in the dynamical ring has also been evidenced by the observation of quadrupolelike collective modes. After the ring formation, the rotation rate, while accelerating, crosses a value where the quadrupole collective mode is at zero frequency, such that any elliptical static anisotropy can excite it resonantly. A very small bubble anisotropy is enough to excite this mode characterized by an elliptic ring shape rotating with a period of approximately 10 s in the direction opposite to the superfluid flow (Fig. 10). This counterpropagating effect is not predicted by a mean-field theory and has been confirmed by resonant spectroscopy of the quadrupole mode during the ring acceleration.<sup>128</sup>

The persistence of superfluidity at such hypersonic velocity raises fundamental questions about the decay of superfluidity in the presence of obstacles, and how superfluidity can be preserved at such speeds: nonlinear effects, the presence of vortices and the dependence on temperature would be particularly interesting to study experimentally and compare with theoretical predictions.<sup>134–138</sup> This hypersonic superflow is not yet a giant vortex, but it is an important step toward this long-sought regime whose transition rotation frequency is not theoretically clearly identified. Moreover, the well-known elementary excitation spectrum for a connected rotating superfluid is strongly modified when the ring appears, and the important discrepancies observed between the experimental results and a mean-field theoretical approach for a quadrupolelike collective mode highlight the need to refine the description of fast rotating superfluids in anharmonic traps.

An alternative way of generating large angular momentum states in rf-dressed adiabatic bubble potentials is to first generate them in a TAAP ring and then reduce the vertical modulation, thus adiabatically transferring the atoms into the bubble.

### C. Trapping in rings with optical potentials

Potentials for ultracold atoms can be formed through the use of focused far-detuned optical beams.<sup>139</sup> Since the potential is directly proportional to the intensity of the optical field, ring-shaped

condensates may be created through the implementation of ring-shaped optical patterns. The most significant advantage in optical dipole ring traps is the insensitivity to the hyperfine state, allowing multicomponent and spinor BECs to be trapped. Additional advantages include the imprinting of superfluid flow, either through phase imprinting or through Raman transitions that can directly transfer angular momentum to the cloud. The advent of spatial light modulator technologies means the optical ring trap has become highly configurable, allowing more complex geometries to be generated.

#### 1. Optical trapping

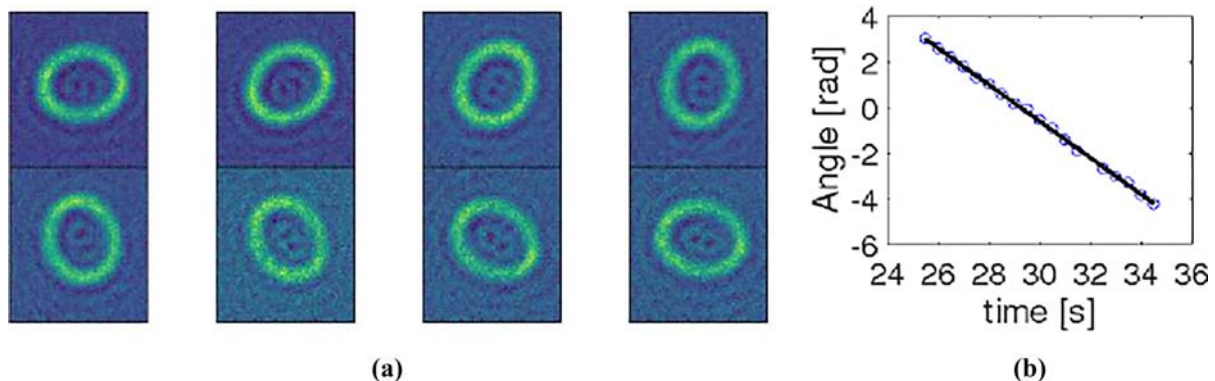
The light-matter interaction can be parameterized through the complex polarizability, where the real part is associated with the dipole trapping potential and the imaginary component results in the absorptive scattering of photons. Trapping cold atoms requires that absorption is minimized to avoid scattering loss of atoms from the trap. Defining  $\Delta = \omega - \omega_0$ , the detuning of the trapping laser from resonance, the scattering loss rate reduces as  $\Delta^{-2}$  while the trapping potential reduces as  $\Delta^{-1}$ . Thus, sufficient detuning of the optical field will result in an optical potential that is approximately conservative. The potential arising for far-detuned dipole trapping light is given by

$$U_{\text{dip}}(\mathbf{r}) = \frac{\pi c^2 \Gamma}{2\omega_0^3 \Delta} I(\mathbf{r}), \quad (11)$$

where  $I(R)$  is the intensity profile of the light and  $\Gamma$  is the transition linewidth. Since the trapping force is determined through the gradient of Eq. (11), a trapping potential requires a nonuniform optical intensity, obtained by shaping and focusing the intensity profile  $I(\mathbf{r})$ . Ring traps, can either be created from attractive (red-detuned) or repulsive (blue-detuned) light, usually by combining the ring shaped intensity profile with a perpendicular light sheet that provides confinement along the propagation direction of the projected ring pattern.

#### 2. Optical ring traps

We begin by looking at some of the optical beam techniques for ring traps that are in use and outline their potential for atomtronic applications.



**FIG. 10.** (a) *In situ* evolution of a quadrupole deformation in the dynamical ring. The elliptical deformation is rotating against the supersonic flow. (b) Time evolution of the orientation of the ellipse major axis.

*a. Laguerre–Gauss beams.* One of the first proposed methods for a ring optical dipole trap was the use of Laguerre–Gaussian (LG) modes having circular symmetry.<sup>140</sup> For far-off-resonance light, these provide the spatial structure for a toroidal trap. An additional advantage of such LG modes is that they also carry orbital angular momentum. With pulses of near-resonant light, the LG modes can be tailored to provide two-photon Raman transitions that transfer exact quanta of circulation to the condensate.

The  $LG_{0N}$  modes are typically generated by phase transformation of a Gaussian  $TEM_{00}$  mode, which transforms the spatial profile of the beam into a donut mode carrying  $N\hbar$  units of orbital angular momentum. A number of methods exist, including spiral phase plates, computer generated holograms, or through the use of phase based spatial light modulators. The toroidal intensity profile of the  $LG_{01}$  mode is given by<sup>141</sup>

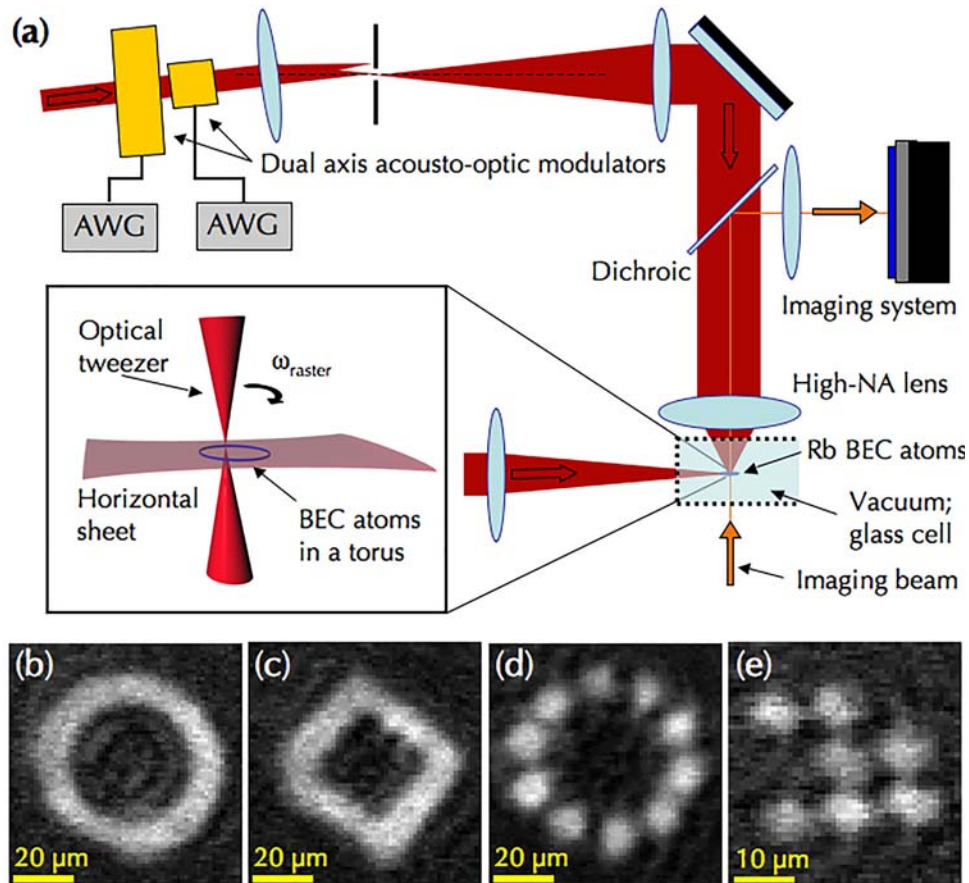
$$I_{LG_{01}}(r) = \frac{4P_{LG_{01}}}{\pi r_0^2} \left(\frac{r}{r_0}\right)^2 e^{-2r^2/r_0^2}, \quad (12)$$

where  $P_{LG}$  is the total laser power in the LG beam and  $r_0$  is the radius at the peak intensity of the LG mode. Correction for imperfections in

the spatial structure, and obtaining sufficient power in higher order modes, is typically a challenge. Ring traps and circulating currents using LG modes have been demonstrated in both single state and multicomponent spinor gases and were early demonstrations of all-optical trapping of BEC in a ring geometry.<sup>67,142,143</sup> To date, they have been used to realize small optical rings for the study of quantized superfluid flows.

*b. Painted optical traps.* An alternative to projecting a ring shaped beam is to build a time-averaged potential with a moving, red-detuned, focused laser beam. By rapidly steering a Gaussian beam in a circular orbit, a ring trap can be generated. This is achieved through the use of two acousto-optic deflectors (AOD) controlling the two axes of the painting beam by driving the deflectors with lists of frequency points that are repeatedly iterated at high speed.<sup>7,32</sup> This approach was used to create the first ring BEC,<sup>7</sup> as shown in Fig. 11.

The advantages of this technique is that it allows adapting the intensity locally to create desired features in the potential landscape and to flatten imperfections due to possible laser inhomogeneity;<sup>28</sup> the available laser power is used in an efficient way as only the relevant trapping locations are illuminated; the painting laser itself can be used



**FIG. 11.** (a) A painted potential system in which a tightly focused rapidly moving red-detuned laser beam paints the desired potential on a horizontal light sheet providing vertical confinement. (b) In-trap absorption images of BECs formed in painted potentials. The technique can create a BEC in any shape that can be drawn on a sheet of paper. Reprinted with permission from Henderson *et al.*, New J. Phys. 11, 043030 (2009). Copyright 2009, Author(s) licensed under a Creative Commons Attribution 3.0 License



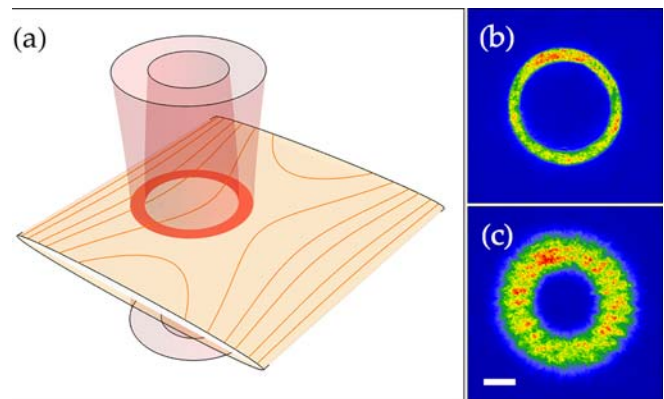
as a stirrer to set the quantum fluid into rotation and demonstrate quantized superfluid flows;<sup>144</sup> the technique also enables more complex geometries. As an example, the atomtronic analog of a Josephson junction has been demonstrated and used to realize a DC atomtronic SQUID.<sup>12</sup> More recently, the dynamic potentials possible with painting were used to show that the atomtronic SQUID exhibits quantum interference.<sup>15</sup>

The painting approach also comes with specific technical constraints that may need to be addressed. The phase of the time-averaged beam loop plays a role on the fine details of the potential, which results in imprinting of the condensate phase, and has to be compensated for.<sup>49</sup> This is particularly relevant for the application of such traps in atom interferometry schemes.

*c. Conical refraction.* A novel approach to generating ring traps has been demonstrated with the use of conical refraction occurring in biaxial crystals. A focused Gaussian beam passing along the optical axis of the crystal transforms, at the focal plane, into one or more concentric rings of light. In the case of a double-ring, the light field encloses a ring of null intensity, called the Poggendorff dark ring.<sup>145</sup> For a blue-detuned laser field, the atoms are trapped between the bright rings. The advantage of this configuration is that it minimizes spontaneous scattering of photons responsible for heating when the laser beam is not very far detuned from resonance. Further advantages include the high conversion efficiency of the incoming Gaussian beam to the ring-trap light field and the access to different ring configurations. The ring diameter is defined by the refractive indices of the biaxial crystal and its length. The width of each ring is given by the focal waist of the focused Gaussian beam. A variation of the ratio of these numbers (e.g., by changing the focal waist) allows for a variation of the resulting light field topology from a single bright ring to a bright ring with a central bright spot and further to bright double rings of increasing diameter. As with LG modes, there are challenges in alignment of the optical beams through the biaxial crystal. On the other hand, the conversion efficiency from a Gaussian TEM<sub>00</sub> mode to the ring pattern can be close to unity. The first results on BECs transferred into a Poggendorff ring have been reported.<sup>145</sup> Ongoing work is directed toward implementing quantum sensors (e.g., Sagnac interferometers) for rings with large diameter and atomtronic SQUIDs for small rings.

*d. Digital micromirror direct projection.* Direct imaging of digital micromirror devices (DMDs) has recently emerged as a powerful tool for the all-optical configuration of BECs.<sup>9,73,146,147</sup> Ring traps can be created by directly projecting the DMD-patterned light onto a vertically confining attractive light-sheet potential,<sup>9,146</sup> similarly to Fig. 12, or onto a vertically oriented accordion lattice.<sup>73</sup> This can be accomplished using a relatively simple optical system, usually consisting of an infinite conjugate pair. Due to the large magnification factors required to reduce the DMD image to the typical 100  $\mu\text{m}$  scale of the BEC, the final element in the imaging system is typically an infinity corrected microscope objective.<sup>9,146</sup> DMDs may also be used in the Fourier plane of the imaging system,<sup>72</sup> where the DMD implements an amplitude-only hologram. A detailed discussion of holographic techniques is beyond the scope of this section, and the reader is referred to more complete reviews of the subject.<sup>148</sup>

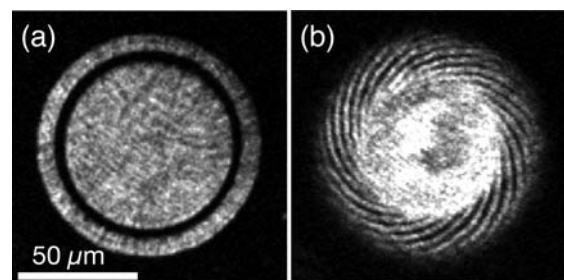
In Fig. 13, direct imaging of a DMD is used to create a ring trap, along with a central phase-uncorrelated reference BEC. By introducing



**Fig. 12.** (a) A typical optical ring trap configuration; the potential is formed at the intersection of the vertically focused ring pattern and horizontal sheet beam. (b) In-trap absorption image of a ring BEC formed in a 164  $\mu\text{m}$  diameter time-averaged optical potential with  $(\omega_r, \omega_z) = 2\pi(50, 140)$  Hz trapping frequencies. (c) Expanded ring after 20 ms time of flight. The scale bar is 50  $\mu\text{m}$ .

a stirring barrier with the DMD, and circulating the barrier around the ring, a 21-quanta persistent current results, corresponding to an angular momentum of  $\sim 132 \hbar$  per atom. The winding number of the current is visualized through interference with the reference central BEC after a short 5 ms time of flight.<sup>149</sup> The DMD technology can also be used to phase imprint an azimuthal light gradient such that angular momentum can be imparted to the atoms<sup>150</sup> and a circulating current created.<sup>151</sup>

*e. Microfabricated optical elements.* An approach combining flexibility, integrability, and scalability can be based on the application of microfabricated optical elements for the generation of complex architectures of dipole traps and guides.<sup>152</sup> It draws its potential from the significant advancement in producing diffraction-limited optical elements with high quality on the micro- and nanometer scale. Lithographic manufacturing techniques can be used to produce many identical systems on one substrate for a scalable configuration.<sup>153</sup> On the other hand, state-of-the-art direct laser writing gives high flexibility in producing unique integrated systems and allows for fast



**Fig. 13.** (a) A ring trap with a diameter of 100  $\mu\text{m}$  is created using a repulsive DMD-patterned potential combined with an attractive horizontal sheet beam. By using the dynamic control of the DMD, a stirring barrier is introduced into the ring and then accelerated through  $90^\circ$ , before being removed from the ring. The resulting persistent current is imaged through interference with the central BEC in a short time of flight, determining a net circulation quanta of  $N = 21$ .

prototyping.<sup>154</sup> Applications range from integrated waveguides and interferometer-type structures<sup>155</sup> to arrays of dipole-traps for quantum information processing<sup>156</sup> and single-atom atomtronics implementations.<sup>157</sup> In combination with DMD-based control of the light field (see Sec. III C 2 d), access to dynamic reconfiguration becomes possible. Integrability is not limited to the generation of light fields for dipole potentials but can be extended to the integration of light sources and detectors or even complex quantum-optical systems such as an entire magneto-optical trap.<sup>152</sup>

### 3. Imperfections in optical traps

Defects in the optical potential will influence the ability to sustain superfluid flow without dissipation, or may introduce unwanted phase perturbations on the condensate if the optical potential is time-varying. We can gain some measure of the significance, and the level of control required for optical traps useful in atomtronics, by considering the density of the BEC in a ring potential. In the Thomas–Fermi limit, with sufficient atom number in the trap, the interaction energies dominate over kinetic energy terms, leading to a simplified GPE equation  $[V(\mathbf{r}) + g|\Psi(\mathbf{r})|^2]\Psi(\mathbf{r}) = \mu\Psi(\mathbf{r})$ , giving the density  $n(\mathbf{r}) = |\Psi(\mathbf{r})|^2 = [\mu - V(\mathbf{r})]/g$ , where  $\mu$  is defined by Eq. (4). The density occupies the spatial profile of the ring trap. In the context of the intensity of the optical potential, assuming a fixed light sheet, the trap depth scales directly scales with the ring optical intensity  $I_0$ , while the chemical potential of the BEC more weakly follows as  $\mu \propto I_0^{1/4}$ . This means that for a typical condensate, the chemical potential is on the order of tens of nanokelvin, and is only weakly effected by the trapping intensity, while large optical trap depths on the order of 1  $\mu\text{K}$  or larger may be easily achieved and utilized. Since the density of the condensate closely follows the optical potential, small perturbations in the optical field can result in significant fluctuations on at the energy scale of the condensate, and thus significant density fluctuations; variations in the optical intensity must typically be limited to less than 1% in order to avoid unwanted perturbations. The precision of the optical projection is thus a key consideration when implementing configurable optical potentials. These aspects, however, also mean that the condensate density provides a very sensitive probe of the optical potential, and the atom density can be used to feedforward corrections to the optical potential.<sup>32</sup>

#### D. Hybrid traps: RF bubble plus light sheet(s)

One can also combine optical potentials and magnetic trapping to produce a hybrid trap and exploit the advantages of each technique for ring trap generation. As mentioned above, optical potentials can achieve large trapping frequencies, while magnetic traps are very smooth due to the macroscopic size of the coils generating them. The bubble geometry described in Secs. III B 1 and III B 3 is particularly suited to create a ring trap: by combining the rf-dressed bubble trap and an optical light sheet as in Sec. III C 2, one can create a toroidal trap. The principle is depicted in Fig. 14: a horizontal light sheet is superimposed with a bubble trap which is rotationally invariant around the vertical direction.<sup>100,158</sup> The light sheet is designed to achieve a strong optical confinement in the vertical direction, and the radial confinement is ensured by the bubble trap itself, made with the same rf-dressed quadrupole trap as in Sec. III B 3. Maximum radial

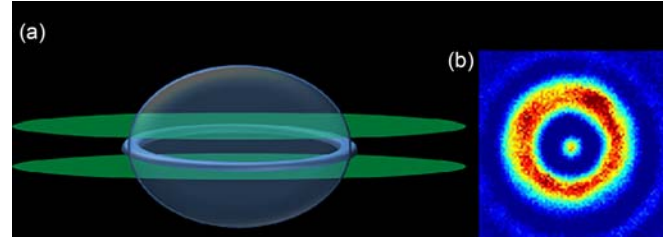


Fig. 14. (a) Principle of the ring trap based on the combination of a magnetic bubble trap and two blue-detuned light sheets. (b) *In situ* image of the ring. The spot at the center is not due to the presence of atoms inside the ring but only to optical diffraction from the ring.

trapping and maximum radius will be attained if the light sheet is located at the equator of the bubble, a situation which also ensures maximum decoupling between the vertical trapping frequency  $\omega_z$  and the radial trapping frequency  $\omega_r$ .

Experimentally, the optical trap is formed between two horizontal light sheets, which are made repulsive by their large blue detuning from the atomic transition. The bubble radius is significantly smaller than the light sheets width and also the vertical Rayleigh length to minimize the azimuthal potential variations. The choice of a small radius also comes with a higher critical temperature and a larger chemical potential, which reduces the relative density fluctuations around the ring due to optical imperfections from residual light scattering of the vacuum glass cell (see Sec. III C 2). One then creates a trapped toroidal degenerate gas of approximately  $10^5$  atoms [Fig. 14(b)]. With further reduction of optical imperfections in the light sheets, one could enter with  $10^4$  atoms the quasi-1D condensate regime,<sup>159</sup> where large-scale correlations and solitons play an essential role in the dynamics.

The gas can be set into rotation by different procedures, using either magnetic or optical means. The first method, used in our experiment in Ref. 160, consists in slightly deforming the bubble trap with an ellipsoidal anisotropy, rotate this magnetic deformation at a given fixed frequency and finally restore the circular symmetry. In a second method (Fig. 15), the rotation is induced by a rotating optical defect<sup>160,161</sup> driven by a dual-axis acousto-optic modulator system as described in Sec. III B 1. Well-controlled circulation could also be imparted by direct optical phase imprinting onto the ring trap.<sup>150</sup>

Above some critical rotation frequency depending on the excitation strength, one observes, after a time-of-flight imaging procedure, a

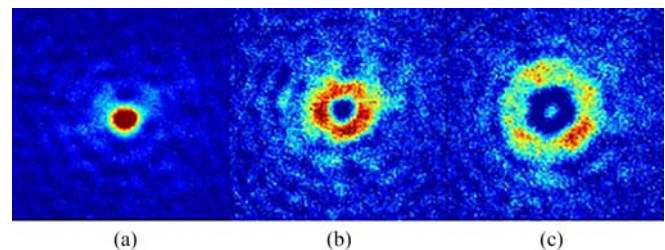


Fig. 15. Time-of-flight images of (a): a nonrotating ring, (b) and (c): rotating rings with different circulations. The rotation is imparted by a rotating 7  $\mu\text{m}$ -waist blue-detuned vertical Gaussian beam.

hole in the atomic distribution. The hole is absent when the ring is nonrotating, and is, thus, evidence for a nonzero circulation of the superfluid in the ring trap (Fig. 15). The hole area grows for increasing rotation rates and shrinks with time when one lets the cloud rotate freely in the trap. In future experiments, optical barriers created by spatial light modulators could be imposed onto the ring and dynamically modulated in height and position. This would create the equivalent of Josephson junctions in superconductors and allow us to simulate models of nonequilibrium quantum systems and emulate new setups in mesoscopic superconductivity.<sup>16,28</sup> This hybrid ring is very promising for the study of 1D superfluid dynamics, for example, shock waves induced by rotation in the presence of a static barrier.<sup>162,163</sup> Increasing again the ring confinement toward the one-dimensional regime with fermionization of the atoms<sup>164</sup> could lead to NOON states more robust against decoherence.<sup>165</sup>

### E. Concluding remarks and outlook

The development of technology for controlling electronic systems and generating complex optical fields is giving ever greater control of ultracold atoms and condensates of atoms. The ring trap remains of particular interest because of the topology, the possibility for self-interference, circuital currents, Sagnac interferometry and so on. In a way, it is its own primitive atomtronic circuit. For optical ring traps, painted optical potentials and digital micromirror devices have demonstrated high level of configurability and dynamic control over the condensate, allowing state-independent trapping, and the ability to introduce junctions, moveable barriers into the atomtronic ring. The ring systems based on RF dressed magnetic traps are also extremely flexible because of the level of electronic control. Atoms can be accelerated and rotated around ultrasmooth waveguides, simply by varying or introducing additional control frequencies with time. The future challenges for the technology, after this development, will be to create particular atomtronic applications and test the limits of technology for creating large scale structures and structures, which possibly have some 3D features. In the future, we will undoubtedly see more control complexity and more hybrid approaches. Where surface interactions are less of a problem, we can also envisage atomtronic circuits based on atom-chip technology, where rings, and complex guided circuits, may be enabled by the design of wire structures and the fields they produce from static and AC currents.

### ACKNOWLEDGMENTS

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### IV. ATOMTRONIC CHIPS AND HYBRID SYSTEMS

**C. Hufnagel, M. Keil, A. Günther, R. Folman, J. Fortagh, R. Dumke**

During the last decade atom chip approaches to quantum technology have become a powerful platform for scalable atomic quantum-optical systems,<sup>104,105,166</sup> with applications ranging from sensor and imaging technologies to quantum processing and memory. Atom chips coupled to solid state-based quantum devices, e.g., superconducting qubits or nitrogen vacancy centers, are thereby paving the way for promising quantum simulation and computation schemes.<sup>167–169</sup> Along this research line, several groups around the world have developed versatile atom chip configurations, which allow trapping of ultracold atomic clouds and degenerate Bose-Einstein condensates (BECs) close to chip surfaces and well-defined manipulation of their internal and external degrees of freedom. Atom chips provide a very relevant technology for the emerging field of atomtronics,<sup>5,30,105,170–173</sup> for which dynamic tunneling barriers are required.<sup>175,179</sup> Such barriers may be formed on atom chips with  $\mu\text{m}$ -scale widths, matching the length scale dictated by the atomic deBroglie wavelength. The atom chip offers the ability to realize guides and traps with virtually arbitrary architecture and a multitude of novel architectures,<sup>176</sup> with a high degree of control over atomic properties, like interactions and spin, enabling new quantum devices.<sup>5,105</sup>

Here we review progress in our groups in Beer Sheva, Tübingen and Singapore on recent developments in atom chip technologies.

#### A. Progress toward on-chip interferometry

The Ben-Gurion University of the Negev (BGU) Atom Chip Group (<http://www.bgu.ac.il/atomchip>) is promoting the idea of atomtronics without light. This entails circuits for atoms based on electric and magnetic traps, guides and tunneling barriers. The vision is for a complete circuit, including particle sources and detection, that makes no use of gravity, e.g., no time of flight for the development of interference fringes. This requirement means that a future technological device could work at any angle relative to gravity.

As a basis for this effort we use the Atom Chip technology developed over the past 20 years.<sup>105,166</sup> An example of a circuit design we plan to implement is a continuous-wave, high-finesse Sagnac interferometer, where the multiple turns enabled by the guiding potential allow miniaturization of the loop while maintaining sensitivity to rotation.<sup>174</sup> In the following we briefly present some of the work that has been done to advance the atomtronics technology.

To begin with, a stable tunneling barrier (in terms of instabilities of tunneling rates) should be no wider than the de Broglie wavelength, which is on the order of  $1\ \mu\text{m}$ . Since the resolution with which we can tailor fields is on the order of the distance from the field source, one must construct the atomic circuit at a distance of no more than a few micrometers from the surface of the chip.<sup>175</sup> At these very small atom-surface distances, several problems must be avoided:

- (1) **Johnson noise.** This is a hindering process as it may cause spin-flips (reducing the trap/guide lifetime), as well as



decoherence. In several papers we have shown ways to combat both effects either by the geometry or by the choice of material.<sup>84,175,177–179</sup> We have also measured Johnson noise and calculated its interplay with phase diffusion caused by atom-atom interactions.<sup>170</sup>

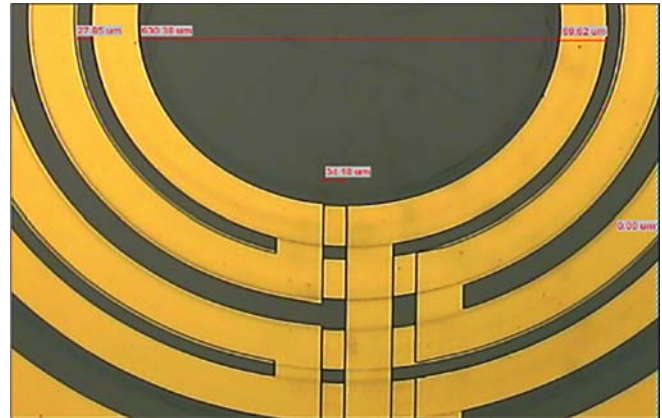
- (2) **Finite size effects.** As the atom-surface distance becomes smaller, so should the current-carrying wire width, or else the magnetic gradients will be severely undermined. Narrow wires require high-resolution fabrication<sup>175</sup> or thin self-assembled conductors such as carbon nanotubes.<sup>179</sup>
- (3) **Casimir-Polder and van der Waals forces.** As the atom-surface distance becomes smaller, the magnetic barrier between the atoms and the surface should be strong enough to avoid tunneling of the atoms to the surface. This has been calculated for submicrometer distances.<sup>175,179</sup>
- (4) **Fragmentation.** Due to electron scattering in the current-carrying wires (e.g., due to rough wire edges), the minimum of the trap, or guide, is not smooth and the atomic ensemble may split and exhibit a nonuniform density along the wire axis. This was studied by us both experimentally and theoretically.<sup>180–182</sup>
- (5) **High aspect ratios.** As the atom-surface distance becomes smaller, the trap or guide exhibits much higher transverse frequencies compared to the longitudinal frequency. This brings about low dimensionality and can cause different problems such as phase fluctuations in a 1D BEC. Alternative wire configurations allow more flexibility for adjusting the trap aspect ratio.

In a proof-of-principle experiment<sup>183</sup> we were able to avoid all the above hindering effects, and showed that spatial coherence could be maintained for at least half a second at an atom-surface distance of just 5  $\mu\text{m}$ .

Another important problem that needs to be overcome is that of atom detection at very small atom-surface distances. At these distances of a few micrometers, the stray light from the nearby surface makes it very hard to achieve a reasonable signal-to-noise ratio for *in situ* detection with typical optical elements. As a solution, and also to avoid on-resonance spin-flips and decoherence, we studied the possibility of off-resonant atom detection with high-Q microdiscs.<sup>184–186</sup>

With the above tools we are now preparing to go forward with our vision for a Sagnac circuit,<sup>174</sup> where as a first stage we have the goal of observing spatial coherence of atoms after one, and then several, turns in a guiding loop. The guiding potential is made in two alternative ways. The first method, using RF potentials, is being led by Thomas Fernholz of the University of Nottingham. It requires multilayer chips (4 layers of currents), which are fabricated at BGU. Two such layers are shown in Fig. 16. The second effort also requires a unique chip. The guiding potential will be based on a repulsive permanent magnet potential in combination with an attractive electric field produced by a charged wire. The first experiments will be pulsed, whereby a BEC will be loaded onto the loop at the beginning of every cycle. Later on we will move toward realizing a continuous-wave version. We will first conduct the experiment in the pulsed mode by loading a thermal cloud, and later on use a 2D MOT as a continuous source.

Finally, let us note that quite a few groups around the world have realized free-space matter wave interferometry. It is now an important challenge to adapt these interferometers to the framework of



**Fig. 16.** A multilayer current-carrying chip produced at BGU for a Sagnac experiment, in co-operation with Thomas Fernholz and Peter Krüger at the University of Nottingham. Concentric gold conducting rings (2  $\mu\text{m}$  thick, upper layer) are 70  $\mu\text{m}$  wide, with intervening gaps of 20–50  $\mu\text{m}$ . The lower layer is also 2  $\mu\text{m}$  thick gold, with a 3  $\mu\text{m}$  thick SU8 insulating layer planarized to better than 0.4  $\mu\text{m}$ . The entire device's outer diameter is 1.37 mm.

atomtronics. Specifically, the BGU Atom Chip group has made significant steps in this direction by realizing, in the last 5 years, several types of novel interferometers which are not based on light. These interferometers are based on the magnetic splitting force (Stern-Gerlach) and they have already enabled the observation of spatial fringes,<sup>187,188</sup> spin population fringes,<sup>189</sup> unique  $T^3$  phase accumulation,<sup>190</sup> clock interferometry,<sup>191,192</sup> and the observation of geometric phase.<sup>193</sup>

## B. Precision sensing

Precise sensors are one of the most important elements in applied and fundamental science. The use of quantum properties in sensing applications promises a new level of sensitivity and accuracy.<sup>194</sup> Using cold atoms on atom chips as probes will enable many interesting applications.

In the laboratories at the University of Tübingen we are working with atom chips that host one or two layers of lithographically implemented wire patterns. They allow the creation of spatially and temporally varying magnetic fields, as used for three-dimensional positioning and manipulation of cold atomic quantum matter.<sup>195</sup> We typically use wire patterns made of gold in room temperature environments<sup>196</sup> and superconducting patterns of Niobium in 4 K and mK surroundings.<sup>197</sup>

With such a “carrier chip” for cold atoms on hand, we established a dual-chip process, where a second chip hosting nanostructured solid state systems is attached on top of the carrier chip.<sup>195</sup> In this way, cold atoms can be efficiently coupled to other quantum systems and hybrid systems can be realized.

We have used this scheme to develop a novel cold-atom scanning probe microscope (CASPM), which uses ultracold atoms and BECs as sensitive probe tips for investigating and imaging nanoscale systems.<sup>198</sup> Similar to an atomic force microscope (AFM), the probe tip is scanned across the surface of interest, while static and dynamical properties of the probe tip are monitored. Evaluating changes within the cold-atom tip density and motion then gives access to basic

interactions and serves as a novel imaging and sensor technique. In contrast to conventional AFMs with their “heavy and rigid” solid state tips, our CASPM uses a dilute gas of atoms, which not only allows for nondestructive measurements, but also for much higher sensitivity to external forces and fields. Inspired by conventional AFMs, we have been able to demonstrate several modes of operation.<sup>199</sup> These include not only a contact mode, where we measure position-dependent losses of the probe tip, but also a dynamic mode, where we initiate a center-of-mass oscillation of the cold-atom tip and monitor the position-dependent changes of the probe tip oscillation frequency.<sup>198</sup> Based on the latter, we have used cold-atom force spectroscopy to unveil anharmonic contributions in near-surface potentials. As in atomic force microscopy, this may be used to reconstruct the surface potentials. Moreover, we have developed a novel operation mode, not accessible to conventional AFMs, where we bring the dilute probe tip into direct overlap with the nano-object of interest. By measuring time-dependent probe tip losses, we have then been able to deduce the underlying van der Waals (Casimir-Polder) interactions.<sup>200,201</sup> We have demonstrated and characterized all different operating modes of CASPM by measuring individual free-standing carbon nanotubes grown on a silicon chip surface. Here we have shown that CASPM extends the force sensitivity of conventional AFMs by several orders of magnitude down to the  $\gamma\text{N}$  regime, and the working distance up to several micrometers.<sup>199</sup> This makes CASPM a powerful tool for investigating fragile nano-objects with ultrahigh force sensitivity.

While first measurements with CASPM suffered from long measurement times, we have just lately extended the microscope by a powerful single atom detection scheme.<sup>202,203</sup> It is based on continuous subsampling of the probe tip via a multiphoton ionization process in conjunction with temporally resolved ion detection and high quantum efficiency. This allows real-time monitoring of the probe tip dynamics and density while losing only few atoms from the probe tip.<sup>203,204</sup> This not only speeds up probe tip oscillation frequency measurements by at least three orders of magnitude,<sup>203</sup> but also enables new applications for CASPM.

In one of these applications we proposed a quantum galvanometer to detect local currents and current noise in nanoscale mechanical quantum devices.<sup>205,206</sup> Measuring the current noise would then give

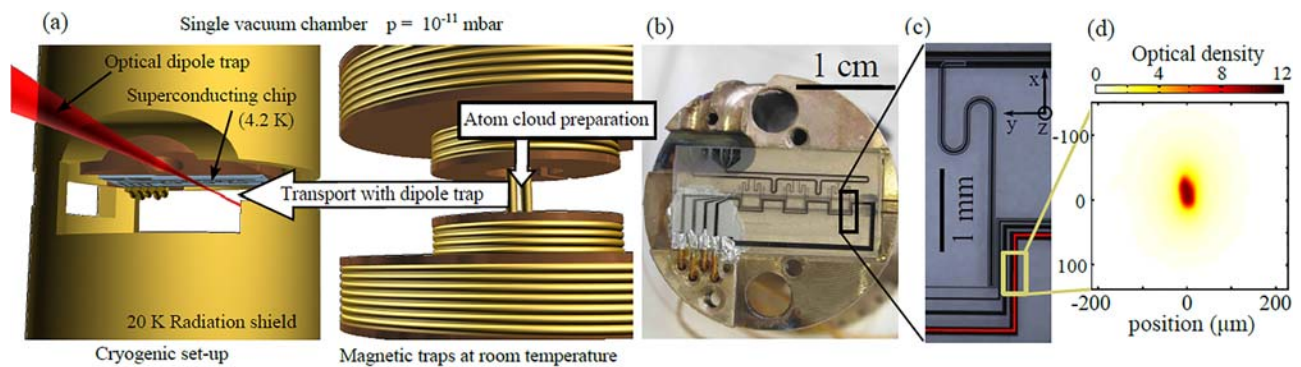
access to the quantum properties of the device. We successfully demonstrated the principal operating scheme of this galvanometer by coherently transferring artificially generated magnetic field fluctuations via a Bose-Einstein condensate onto an atom laser and investigating its single-atom statistics.<sup>207,208</sup> Employing second-order correlation analysis, we could not only extract the microwave power spectral density (current noise spectrum) but also the noise correlations within the bandwidth of the BEC, which will give access to the quantum noise properties of the current source. This will extend CASPM to a promising quantum sensor, not only for detecting local forces and force gradients, but also for currents as well as electric and magnetic fields (AC and DC), including their specific noise spectra.

### C. Cryogenic atom chips and hybrid quantum systems

Atom chips made from superconducting circuits offer certain advantages over normal metal devices. The coherence properties of trapped atoms are improved by orders of magnitude due to reduction of magnetic noise coming from the surface of the chip. Moreover, superconductors can be operated in the mixed state, where vortices can be used to generate self-sufficient atom traps. In addition to that, working in cryogenic environments offers the possibility to interface atoms with solid state devices to form hybrid quantum systems.<sup>84</sup>

Besides atom chip experiments in room-temperature environments, the group in Tübingen also operates superconducting atom chips with trapped BECs of rubidium atoms.<sup>197,209,210,214</sup> As shown in Fig. 17, condensates are routinely transferred into coplanar cavity structures<sup>211</sup> and the measured coherence time between hyperfine ground state superpositions reaches several seconds. Microwave dressing is used to suppress the differential shift of state pairs with the “double-magic point” being the optimum working point for quantum memories.<sup>212</sup> We have successfully demonstrated coherent coupling of a hyperfine state pair through a driven superconducting coplanar microwave cavity,<sup>213</sup> which paves the way for future cavity-based quantum gate operations.

In addition to manipulating ground-state atoms we have successfully implemented two-photon Rydberg excitation in a cryogenic environment near the superconducting chip.<sup>215</sup> We have developed techniques for optical detection of Rydberg populations and



**FIG. 17.** Superconducting atom chip at the University of Tübingen. (a) *In vacuo* trap setup with electromagnets for cold atom preparation (right) and cryostat with a superconducting chip at 4.2 K (left). (b) Photograph of the chip holder and the superconducting atom chip. (c) The chip features superconducting wire components for magnetic trapping and positioning of atomic clouds and a coplanar microwave cavity. (d) Bose-Einstein condensate of  $3 \times 10^5$   $^{87}\text{Rb}$  atoms in a 15 ms time-of-flight image, released from the superconducting atom chip. Reprinted with permission from Bernon *et al.*, Nat. Commun. 4, 2380 (2013). Copyright 2013, Nature Publishing Group.



coherences<sup>216</sup> and measured the increased lifetime of Rydberg states in cryogenic environments.<sup>215</sup> In preparation for interfacing Rydberg atoms with superconducting circuits, we have obtained high-resolution spectra of rubidium Rydberg states in a field-free vapor cell as reference,<sup>217</sup> and in precisely controlled electrostatic fields<sup>218</sup> near surfaces at room and cryogenic temperatures. These studies add to our understanding of electrostatic fields of surface adsorbates that build up during experiments with cold atoms at chip surfaces.<sup>88,219,220</sup> Based on the measured data, quasi-classical quantum defect theory,<sup>221</sup> Stark-map calculations,<sup>218</sup> suitable dressing techniques,<sup>212,222</sup> and numerical methods developed for simulating quantum operations in the presence of thermal cavity photons,<sup>89</sup> we are currently focusing on the coherent manipulation of Rydberg atoms and quantum computation schemes in the presence of inhomogeneous fields at the surface of superconducting coplanar cavities.

The realization of hybrid quantum systems based on atoms and superconducting qubits requires truly cold temperatures in the 10 mK range, as dictated by the otherwise fast decoherence of the superconducting qubit. The great advantage is that at this temperature the number of microwave photons in the cavities that mediate the interaction between the solid state and the atomic system is near zero. The price to pay is a highly complex experimental system combining cold-atom technologies with a <sup>3</sup>He/<sup>4</sup>He dilution refrigerator.<sup>223,224</sup> Our dilution refrigerator consists of several temperature-shielded volumes (stages), of which we use the 6 K-stage and the 1 K-stage for cold atom experiments. The 1 K-stage includes a cold plate with a nominal base temperature of 25 mK.<sup>209</sup> We routinely operate a magneto-optical trap at the 6 K-stage, from which we transport magnetically trapped, ultracold rubidium clouds at 100  $\mu$ K to the 1 K-stage. The 1 K-stage has a sufficiently large volume (several liters) to accommodate microwave cavities, such as coplanar waveguide cavities, and has convenient optical access for optical traps and laser beams for spectroscopic measurements. This experimental setup is currently being extended for studying the fully quantum regime of cold-atom superconductor hybrid systems.

In the Singapore group we are working in two directions. One is the exploration of superconducting atom chips using high-temperature superconductors and another is the development of coherent interfaces between superconducting circuits and ultracold atoms.

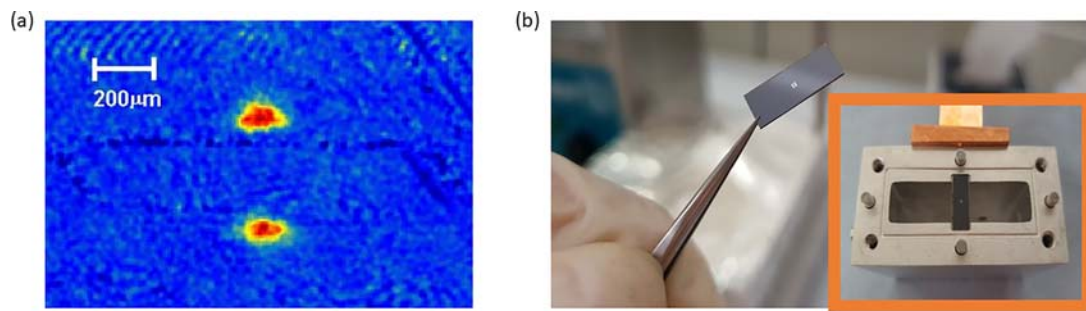
High temperature superconductors have various distinct properties when implemented as atom chips. First of all, the technical

demands are lower due to the higher working temperatures, which can be reached with liquid nitrogen instead of liquid helium. Moreover, high temperature superconductors are type-II superconductors and allow the storage of magnetic fields in the remanent state. We have shown experimentally and in simulations, that these trapped fields can be used to generate novel traps for ultracold atoms.<sup>225,226</sup> Ramping a magnetic field perpendicular to a planar structure of YBCO we were able to generate various magnetic traps for cold atoms [see Fig. 18(a)].<sup>227</sup> These traps can be generated either by using external magnetic fields together with vortices or in a completely self-sufficient way, where the trap is solely created by vortices. In the latter case, low noise potentials can be generated, as there is no technical noise coming from external power sources and the noise coming from the movement of vortices is expected to be an order of magnitude less than Johnson noise in normal conductors.<sup>84</sup>

Another property of vortices in superconducting thin films is that their distribution can be manipulated with light. Heating parts of the superconductor will result in a force on the vortices, which shifts the position of the vortices and consequently changes the vortex distribution. We have used this effect to generate various trap patterns with a thin square of superconducting YBCO, using light patterns generated by a spacial light modulator.<sup>228</sup> The advantage of this technique is that multiple trap geometries can be generated with the same chip architecture in-situ, without the need of changing the chip and breaking the vacuum of the ultrahigh vacuum chamber.

Aside from using superconducting chips exclusively to manipulate cold atoms, we are also working on interfaces between cold atoms and superconducting qubits fabricated on the superconducting chip. What we envision here is the coherent transfer of quantum states between cold atoms and qubits made of superconducting integrated circuits. These hybrid systems will have many application, like the transduction of quantum states between the microwave and optical regime or the creation of universal quantum computing devices.

As mentioned before in this article, the practical implementation of a hybrid atom-superconducting qubit system is technically challenging. In Singapore we decided to bring cold atoms inside the dilution refrigerator by magnetically transporting them from a room temperature vacuum chamber directly to the mK stage of the refrigerator. With this technique we are able to bring clouds of  $5 \times 10^8$  <sup>87</sup>Rb



**Fig. 18.** (a) Image of a thermal atomic cloud of  $10^3$  <sup>87</sup>Rb trapped in a self-sufficient quadrupole magnetic trap generated by vortices on a  $1 \times 1$  mm<sup>2</sup> square of superconducting YBCO (National University of Singapore). The lower part of the image shows a mirror image of the cloud, caused by the reflection of the imaging beam from the chip surface. Reprinted with permission from Siercke *et al.*, Phys. Rev. A **85**, 041403 (2012). Copyright 2012, American Physical Society. (b) Setup for a hybrid quantum system of atoms and superconductors. The image shows a transmon qubit fabricated on a silicon substrate. Inset: the transmon installed in a 3D microwave cavity. The cavity allows to read-out and manipulate the transmon with RF radiation. In the future, we plan to bring atoms into the cavity to form a hybrid quantum system.

atoms close to the mK stage, at a base temperature of 70 mK.<sup>224</sup> Trapped inside the mK stage, the atomic cloud exhibits an exceptional lifetime of 13 minutes, which is a promising starting point for future experiments.

In order to couple atoms and superconducting circuits a few scenarios are possible, which can be categorized in indirect and direct coupling. Also, the state of the atoms, i.e., ground state or highly excited (Rydberg), will have a significant influence on the experimental parameters. When indirectly coupled, the qubit and atom are individually coupled to a resonator, which mediates the interaction. In this case the coupling of the resonator to the qubit is easily implemented and can reach the strong coupling regime. Coupling ground state atoms to a planar resonator is an ambitious task. It was shown that the coupling strength of a single atom is only 40 Hz at a resonator-atom distance of 1  $\mu\text{m}$ .<sup>229</sup> In order to reach strong coupling one consequently needs to collectively couple an ensemble of  $10^6$  atoms to the waveguide, which is experimentally challenging. Using Rydberg states can considerably relax these requirements. We have shown that for Rydberg states strong coupling can be achieved with even a single atom.<sup>90,230</sup> The strong coupling can even be reached with atom-resonator distances of tens of micrometers, when using the fringe field of the capacitive part of the resonator to couple the atom.

When using Rydberg atoms, even directly coupling of atoms to charge qubits can be realized. A neutral atom placed inside the gate capacitor of a charge qubit acts as a dielectric medium and affects the gate capacitance, resulting in a modulation of the charge-qubit energy bands. Moreover, the local quasi-static electric field strongly depends on the charge-qubit state, leading to different DC Stark shifts of atomic-qubit states. We have shown that in such a setup quantum states can be transferred between the two qubits and CNOT and Hadamard gates can be realized.<sup>231</sup> Schemes for Rydberg atoms interacting with flux qubits have been theoretically proposed to realize quantum memories.<sup>232</sup>

We think that we now have the tools at hand to interface cold atoms with superconducting circuits. In the near future we would like to first couple atoms to 3D transmons, see Fig. 18(b). For this we designed and tested superconducting 3D cavities that have free space access for the transport and optical manipulation of cold atoms. First experiments to transport atoms inside the cavity are under way. At the same time we are developing our own fabrication for superconducting qubits. First chips have already been fabricated and tested. With both systems at hand we can then go forward to build hybrid systems of cold atoms and superconducting circuits.

## D. Concluding remarks and outlook

In this review we have described applications of atom chips in atomtronics, precision sensing and quantum information. We illustrated the state of the art in these topics and touched upon future prospects and utilizations. In this zoomed-in view, we omitted many other excellent activities in the field, due only to unavoidable space limitations. Here, we would like to bring up other achievements that will shape the future of the atom chip platform.

Most of the experimental studies described above used bosonic rubidium atoms. In fact, many other species, like fermions, molecules and ions are used in atom chips.<sup>105</sup> Fermions are another one of the fundamental building blocks of matter and therefore highly interesting objects to study, including low dimensional physics, the interaction of fermions with different species, or spin physics.<sup>233</sup>

Molecules, as the bridge between physics and chemistry, are an additional compelling candidate for many studies. Implementations range from fundamental science, like the measurement of the electric dipole moment and parity violation, to applied science in quantum processing. A “Lab on a Chip” for molecules is thus a sought-after goal. Recently, the trapping of simple molecules on microchips was realized,<sup>234,235</sup> opening the way for many interesting investigations.

Trapped ions are one of the most promising candidates for practical quantum computing. In order to control and measure a large number of ions it will be necessary to fabricate surface-electrode traps on miniaturized microchips. The development and integration of these chips is currently ongoing and will be a major part in the future development of scalable quantum computer architectures with ions.<sup>236</sup>

Using the wave nature of atoms, atom chips will in future be used as precise sensors for material research and fundamental science. So-called “quantum gas microscopes” have been developed for room-<sup>181</sup> and cryogenic<sup>237</sup>-temperature environments and are ready to be used in the nontrivial studies of unique materials. At the same time, matter waves are being employed for precision measurements in atomic interferometers. By analogy to the optical interferometer, the splitting and recombination of matter waves on atom chips are, for instance, being used to test theories in quantum thermodynamics,<sup>238</sup> quantum many-body physics,<sup>239</sup> and find applications in gravitational sensing.<sup>240</sup>

Intimately connected with precision sensing is the field of fundamental science. Many studies will be possible with atom chips, including tests of the Weak Equivalence Principle,<sup>241</sup> interactions of antihydrogen with matter and gravity,<sup>242</sup> non-Newtonian gravity, and the search for a fifth fundamental force.

All these examples show that atom chip technology has a bright future ahead. Combined with further integration and miniaturization, atom chips will play a role in many areas, both in fundamental research, as well as practical measurements.

## ACKNOWLEDGMENTS

All groups are very thankful to their colleagues who have been working with them on the experiments and their interpretation.

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## V. QUENCH DYNAMICS OF INTEGRABLE MANY-BODY SYSTEMS

**N. Andrei, C. Rylands**

The study of nonequilibrium quantum physics is currently at the intellectual forefront of condensed matter physics. One-dimensional systems in particular provide an exciting arena where over the last decade significant advances in experimental techniques have allowed

very precise study of an array of nonequilibrium phenomena and where a number of powerful theoretical tools were developed to describe these phenomena. Here we give a brief account of a few systems that are described by one dimensional integrable Hamiltonians, the Lieb-Liniger model and the Heisenberg chain and how integrability gives access to the study of some of their local and global nonequilibrium properties.

While the principles of equilibrium statistical mechanics are well understood and form the basis to describe a variety of phenomena, there is no corresponding framework for the nonequilibrium dynamics, although efforts to fully understand the underlying principles extend back to Boltzmann and beyond. Solving particular models numerically or analytically and comparing to experiments may illuminate bits of the puzzle.

Here, is an extended version of talks given by the first author at Atomtronics 2019 at Benasque where some aspects of the questions were discussed. It is based on a review article<sup>243</sup> written with Colin Rylands and builds on work carried out with several collaborators: Deepak Iyer, Garry Goldstein, Wenshuo Liu, Adrian Culver, Huijie Guan and Roshan Tourani to whom we are very grateful for many enlightening and useful discussions.

## A. Quench dynamics

A convenient protocol to observe a system out of equilibrium is to prepare it in some initial state  $|\Phi_i\rangle$ , typically an eigenstate of an initial Hamiltonian  $H_i$ , and then allow it to evolve in time using another Hamiltonian,  $H$  for which  $|\Phi_i\rangle$  is not an eigenstate.<sup>244–246</sup> One then follows the correlations of local observables,

$$\langle \Phi_i | e^{iHt} \{ \mathcal{O}_1(x_1) \mathcal{O}_2(x_2) \dots \} e^{-iHt} | \Phi_i \rangle \quad (13)$$

as they evolve. One may be interested to know what new properties characterize the system, whether a dynamical phase transition occurs at some point in time<sup>247</sup> or how its entanglements evolve. A particularly important question that arises in this context is whether the system thermalizes. Namely, can the system act as a bath to a small subsystem, here the small subsystem is the segment that contains the local operators  $\mathcal{O}_j(x_j)$ . In the long time limit (to thermalize it is necessary that  $vt \gg L$  where  $L$  is the size of the system, and  $v$  a typical velocity) one needs to show

$$\lim_{t \rightarrow \infty} \langle \Phi_i | e^{iHt} \{ \mathcal{O}_1(x_1) \mathcal{O}_2(x_2) \dots \} e^{-iHt} | \Phi_i \rangle = \text{Tr } e^{-\beta H} \{ \mathcal{O}_1(x_1) \mathcal{O}_2(x_2) \dots \} / Z \quad (14)$$

with the final inverse temperature  $\beta$  determined by the initial energy,  $E_0 = \langle \Phi_i | H | \Phi_i \rangle = \text{Tr } e^{-\beta H} H / Z$ .

Also global properties are of interest. These are commonly studied via the Loschmidt amplitude (LA), the overlap between the initial state with its time evolved self, conveniently expressed using a complete set of energy eigenstates,  $|n\rangle$ :

$$\mathcal{G}(t) = \langle \Phi_i | e^{-iHt} | \Phi_i \rangle = \sum_n |\langle n | \Phi_i \rangle|^2 e^{-iE_n t} \quad (15)$$

and its Fourier transform,

$$\mathcal{P}(W) = \int_{-\infty}^{\infty} \frac{dt}{2\pi} e^{iWt} \mathcal{G}(t) = \sum_n \delta(W - E_n) |\langle n | \Phi_i \rangle|^2, \quad (16)$$

which measures the work distribution done during the quench.<sup>248</sup>

## B. Evolution under integrable Hamiltonians

We shall consider evolutions effectuated by post-quench Hamiltonians that are integrable, namely Hamiltonians admitting a complete set of eigenstates  $|n\rangle$  and eigen-energies  $E_n$  given by the Bethe ansatz. The ability to obtain these follows from the existence of an infinite set of local charges,  $\{Q_n, n = 1 \dots \infty\}$ , that commute with the Hamiltonian and constrain the time evolution leading to a generalized Gibbs ensemble  $e^{-\sum_n \beta_n Q_n}$  with the final inverse temperatures  $\beta_n$  determined by the initial values  $q_n^0 = \langle \Phi_i | Q_n | \Phi_i \rangle$ .<sup>249</sup>

Thus some features of integrable time evolution are nongeneric. It turns out however that many features observed in integrable models can also be observed when integrability is broken. An example is the “dynamical fermionization” of repulsively interacting bosons in the integrable Lieb-Liniger Hamiltonian, discussed below. We showed this feature can be also observed in the bose-Hubbard model, the lattice version of the Lieb-Liniger model, which is not integrable.<sup>250</sup> Further, many systems, in particular ultracold atom systems, are actually described by integrable Hamiltonians and can therefore be studied as such. Here we discuss two of them.

### 1. The Lieb-Liniger model

The model describes systems of ultracold gases of neutral bosonic atoms moving in one dimensional traps and interacting with each other via a local density interaction of strength  $c$  which can be repulsive  $c > 0$  or attractive  $c < 0$ . Aside from being an excellent description of the experimental system, it is one of the simpler Hamiltonians for which there exists an exact solution via Bethe ansatz. The Lieb-Liniger Hamiltonian reads

$$H = - \int dx \Psi^\dagger(x) \frac{\partial^2}{\partial x^2} \Psi(x) + c \int dx \Psi^\dagger(x) \Psi(x) \Psi^\dagger(x) \Psi(x) \quad (17)$$

(setting  $\hbar = 1$ ). Here  $\Psi^\dagger(x)$ ,  $\Psi(x)$  create and annihilate bosons of mass  $m$ . The exact  $N$ -particle eigenstate is given by<sup>251,252</sup>

$$| \{k_j\} \rangle = \int d^N x \prod_{i,j=1}^N [\theta(x_i - x_j) + s(k_i, k_j) \theta(x_j - x_i)] \times \prod_{l=1}^N e^{ik_l x_l} \Psi^\dagger(x_l) | 0 \rangle. \quad (18)$$

Here  $s(k_i, k_j) = (k_i - k_j + ic/k_i - k_j + ic) = e^{i\varphi(k_i - k_j)}$  is the two particle scattering matrix,  $\varphi(k) = 2\arctan(k/c)$  is the phase shift. The single particle momenta  $k_j$  are unrestricted in the infinite volume limit while with periodic boundary condition on a line segment  $L$  they must satisfy the Bethe ansatz equations:  $k_i L = \sum_{j=1}^N \varphi(k_i - k_j) + 2\pi n_i$ , with the integers  $n_i$  being the quantum numbers of the state. The single particle momenta are related to the conserved charges by  $q_n = \sum_{j=1}^N k_j^n$ , in particular the energy is given by,  $q_2 = E = \sum_{j=1}^N k_j^2$ .

This set of eigenstates allows the study of time evolution through the partition of the unity,  $\mathbb{1}_N = \sum_{k_1, \dots, k_N} (| \{k_j\} \rangle \langle \{k_j\} | / \mathcal{N}(\{k_j\}))$ .



Here  $\mathcal{N}(\{k\}) = \det[\delta_{jk}(L + \sum_{l=1}^N \varphi'(k_j - k_l)) - \varphi'(k_j - k_k)]$  is a normalization factor.

In terms of the partition identity, the time evolved wavefunction is given by,

$$\begin{aligned} |\Phi(t)\rangle &= e^{-iHt} |\Phi_i\rangle \\ &= \sum_{k_1, \dots, k_N} e^{-it(\sum_{j=1}^N k_j^2)} |\{k_j\}\rangle \langle \{k_j\} | \Phi_i \rangle \end{aligned} \quad (19)$$

with the initial state  $|\Phi_i\rangle$  encoded in the overlaps  $\langle \{k_j\} | \Phi_i \rangle$ . These overlaps have been studied by many groups, e.g.<sup>253</sup> and are typically very difficult to calculate. Once these overlaps are known they can be put in exponential form and combined with matrix elements of a given operator to yield a quench action which is typically evaluated in the saddle point approximation.<sup>254</sup>

**Beyond overlaps:** One may get around the difficulty of computing overlaps by choosing an alternate form of the partition identity obtained by exchanging the ordering in momentum space for ordering in coordinate space leading to the Yudson representation of the partition of the unity,<sup>255,256</sup> or equivalently choosing appropriate trajectories for integrating over the momenta, see<sup>250</sup>

$$1_N = \sum_{k_1, \dots, k_N} \frac{|\{k_j\}\rangle \langle \{k_j\}|}{\mathcal{N}(\{k_j\})}. \quad (20)$$

Here we have introduced the notation  $|\{k_j\}\rangle$  (notice the parenthesis replacing the ket) to denote the Yudson state,

$$|\{k_j\}\rangle = \int d^N x \theta(\vec{x}) \prod_l e^{ik_l x_l} \Psi^\dagger(x_l) |0\rangle \quad (21)$$

with  $\theta(\vec{x})$  denoting a Heaviside function which is nonzero only for the ordering  $x_1 > x_2 > \dots > x_N$ . The Yudson state is simpler to work with than the full eigenstates of the model and its overlaps with the initial state can be readily calculated, particularly if the initial state is ordered in coordinate space.

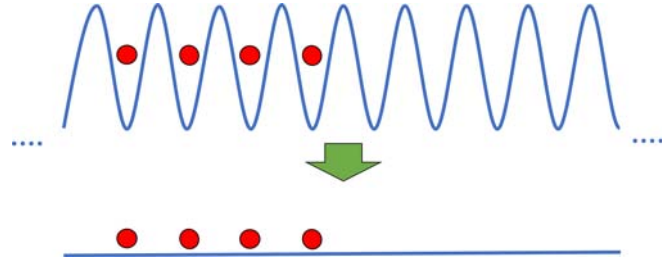
**The domain wall initial state:** As an example we consider an initial state in the form of a domain wall and quench it with  $c > 0$  Lieb-Liniger Hamiltonian. Its time evolution can be studied analytically and several interesting phenomena will be shown to emerge:

- Nonequilibrium Steady state (NESS)
- RG flow in time
- Evolution along space-time rays
- Hanbury Brown-Twiss effect
- Dynamical fermionization

The initial state, as depicted in Fig. 19, consists  $N$  cold atom bosons held in a very deep optical lattice of length  $L$  with  $N, L \rightarrow \infty$  and  $\delta = L/N$  held fixed. The lattice site  $\bar{x}_j = j\delta$ ,  $j = -\infty, \dots, -1, 0, 1, \dots, +\infty$ , are filled with one boson per site in the left half of the lattice:  $j = -\infty$  to  $j = 0$ , and none in the half to the right,

$$|\Phi_i\rangle = \int d^N x \prod_{j=-\infty}^0 \left[ \frac{\omega}{2\pi} \right]^{\frac{1}{4}} e^{-\omega/4(x_j - \bar{x}_j)^2} \Psi^\dagger(x_j) |0\rangle. \quad (22)$$

The quench consists of suddenly releasing the trap and allowing the bosons to interact and evolve under the Lieb-Liniger Hamiltonian.



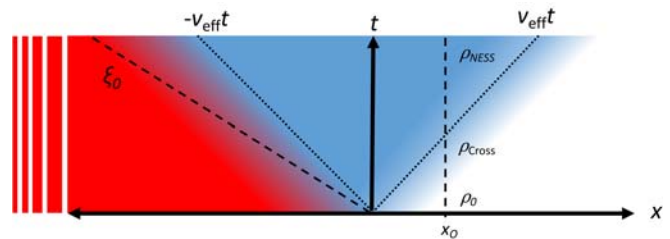
**Fig. 19.** The domain wall initial state: A cold atom gas is held in the left part of a deep optical lattice, extending from  $j = -\infty$  to  $j = 0$ . This is then removed and the gas is allowed to expand. The system is open—its size is large compared to time of evolution.

Time evolving the system and using the Yudson representation we find,

$$|\Phi_i(t)\rangle = \left[ \frac{8\pi}{\omega} \right]^{\frac{N}{4}} \sum_{k_1, \dots, k_N} \frac{e^{-\sum_{j=1}^N [1/\omega(1+i\omega t)k_j^2 + ik_j \bar{x}_j]}}{\mathcal{N}(\{k_j\})} |\{k_j\}\rangle. \quad (23)$$

When the lattice is removed the gas expands and the particle density will become nonzero between the lattice sites and also to the right of the domain wall. In the vicinity of the domain wall particles will begin to vacate the left hand side of the system and populate the right hand side, see Fig. 20. The effects of this quench can only be felt within a “light-cone” centered at the edge and determined by a finite effective velocity,  $v_{\text{eff}}$  which depends upon  $\omega$ . On the right,  $x \gg v_{\text{eff}} t$  the density will remain zero while to the left,  $x \ll -v_{\text{eff}} t$ , the average density will remain  $1/\delta$  - the effects of the quench are still felt as the initially confined bosons will expand and begin to interact with each other.

We first examine the local portion of the quench around the domain wall. Since to the left there is an infinite particle reservoir and to the right an infinite particle drain the system will never equilibrate, however at long times a nonequilibrium steady state (NESS) consisting of a left to right particle current is established. This can be investigated by computing the expectation value of the density  $\rho(x, t) = \langle \Phi_i(t) | \Psi^\dagger(x) \Psi(x) | \Phi_i(t) \rangle$ . Utilizing the known formulae for the matrix elements of the density operator with Bethe eigenstates<sup>257</sup> this can be calculated exactly. To the right of the domain wall, at long times and to leading order in  $1/c\delta$  three regions emerge:<sup>256</sup>



**Fig. 20.** At long times, a nonequilibrium steady state (NESS) is established as depicted on the right. Measuring the density at  $x = x_0$  one will see the initial density  $\rho_0$  change to the crossover regime  $\rho_{\text{cross}}$  at intermediate times ending up as time and space independent value  $\rho_{\text{NESS}}$  which encodes the interaction and the initial quench. Reprinted with permission from Rylands and Andrei, Annu. Rev. Condens. Matter Phys. 11, 147–168 (2020). Copyright 2020, Annual Reviews.

$$\rho(x, t) = \begin{cases} \rho_{\text{NESS}} = \frac{1}{2\delta} - \frac{4\pi}{c\delta^2}, & \frac{1}{\sqrt{\omega}} \ll x \ll v^{\text{eff}} t \\ \rho_{\text{Cross}}(x) = \frac{1}{\delta} f + \frac{16}{\pi c \delta^2} \left[ e^{-(x^2/\sigma)} \frac{x\sqrt{\pi}}{\sqrt{\sigma}} f - \frac{1}{2} e^{-2(x^2/\sigma)} + \frac{\pi}{2} f(1-f) \right], & x \sim v^{\text{eff}} t \\ \rho_0 = 0, & x \gg v^{\text{eff}} t, \end{cases} \quad (24)$$

where  $f = f(x, t) = \frac{1}{2} \text{erfc}(x/\sqrt{\sigma(t)})$  and  $\sigma(t) = t^2 \omega/2 + 2/\omega$ . Far to the right  $x \gg v^{\text{eff}} t$  we see that the density vanishes while closer to the light-cone some complicated crossover behavior occurs. Since the model is Galilean rather than Lorentz invariant the light-cone is not sharp giving instead this crossover regime. Most interesting is the region deep inside the light-cone in which the density becomes independent of  $x, t$ , signifying the appearance of the NESS, Nonequilibrium Steady State. We note that the particle density in this regime,  $\rho_{\text{NESS}} = 1/2\delta - 4\pi/c\delta^2$ , is reduced as compared to the equilibrium value,  $\rho_{\text{EQL}} = 1/2\delta$ , the value a closed system would have reached after a quench from a domain wall state. This is a nonequilibrium effect of an open system which allows the bosons to expand further to the right in response to the repulsive interactions among the bosons. It follows from the order of limits with the size of the system  $L$  satisfying  $L \gg vt$ , to be contrasted with the behavior in a closed system, with the opposite order of limits. Within this region all local properties of the system can be calculated by taking the expectation value with respect to this NESS,  $\langle \mathcal{O}(x, t) \rangle = \langle \Psi_{\text{NESS}} | \mathcal{O} | \Psi_{\text{NESS}} \rangle$  where  $|\Psi_{\text{NESS}}\rangle$  can be determined by taking the appropriate limit of (23).

On the left portion of the lattice  $x \ll -v^{\text{eff}} t$  we are outside the light-cone, the system is unaffected by the domain wall portion of the quench and the lattice translational invariance is restored. At long times the density within this region is

$$\rho(x, t) = \frac{1}{\delta} \left[ 1 + \sum_{s=1}^{\infty} e^{-\sigma(t)(\pi^2 s^2 / \delta^2)} \cos\left(\frac{2\pi s x}{\delta}\right) \right], \quad (25)$$

which describes small oscillation about a uniform density of  $1/\delta$ .

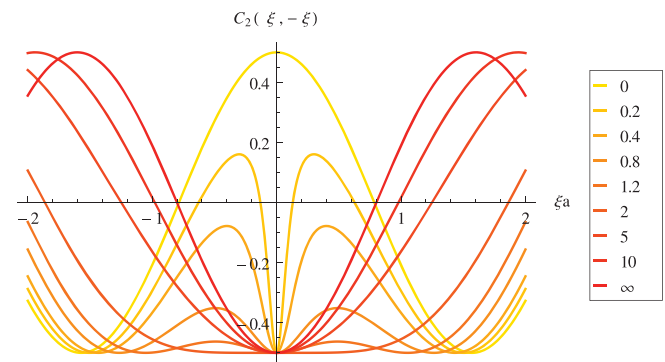
This result coincides with what one would expect for a quench from a lattice initial state of the Tonks-Girardeau (TG) gas, the  $c \rightarrow \infty$  limit of the LL model. To understand this one should go beyond the density and compute the normalized noise correlation function  $C_2(x, x') = \rho_2(x, x', t) / \rho(x, t) \rho(x', t) - 1$  where

$$\rho_2(x, x', t) = \langle \Phi_i(t) | \Psi^\dagger(x) \Psi(x) \Psi^\dagger(x') \Psi(x') | \Phi_i(t) \rangle. \quad (26)$$

This correlation function is related to the Hanbury Brown-Twiss effect and will detect the nature of the interactions between particles, a peak indicating bosons while a dip indicates fermions.<sup>258,259</sup> Computing the noise correlation function  $\rho_2(x, -x, t)$  by inserting two copies of the identity and evaluating the integrals at long time by saddle point method<sup>260</sup> one finds it becomes a function only of the ray variables  $\xi = x/t$  (measured with respect to  $\xi_0 = x_0/t$  see Fig. 20). For sufficiently long times  $\xi \sim 0$  a distinct fermionic dip is seen for arbitrary  $c > 0$  while  $c = 0$  shows a bosonic peak, the turn over to the dip occurring on the time scale,  $t \sim c^{-2}$ , see Fig. 21. This turn over results from an increase in time of the effective coupling constant  $c$  - starting from any initial repulsive value it will flow to strong coupling in the long

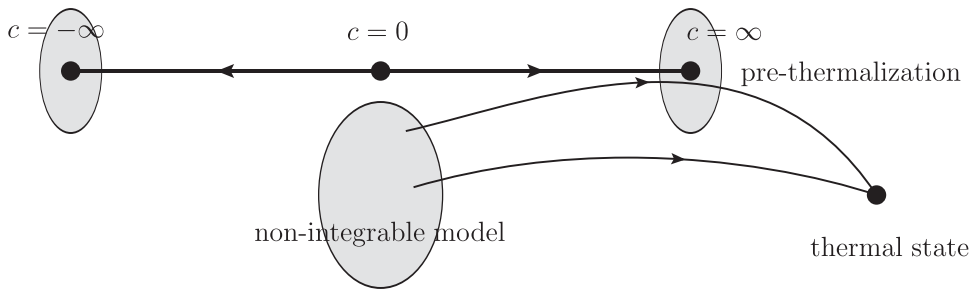
time limit.<sup>261,262</sup> This follows elegantly from the Yudson representation of the time evolving wave function:<sup>260</sup> rewriting the dynamic factor in Eq. (18),  $\theta(x_i - x_j) + s(k_i, k_j) \theta(x_j - x_i)$ , as  $k_i - k_j - ic \text{sgn}(x_i - x_j)/k_i - k_j - ic$ , we note it tends to  $\text{sgn}(x_i - x_j)$  upon rescaling  $k_j^2 t \rightarrow k_j^2$ . Therefore, the product of bosonic fields with the dynamic factors,  $\prod_{i < j} \text{sign}(x_i - x_j) \prod_j \Psi^\dagger(x_j)$ , behaves fermionically. The physical argument underlying the mathematical manipulations is also simple. In the long time limit only the slow bosons remain around  $x, x'$  in the noise correlation function  $\rho_2(x, x', t)$  and they interact via the effective S-matrix  $S^{ij} \rightarrow -1$ . The system in the long time limit will then behave as if it consisted of noninteracting fermions. This dynamical fermionization, the development of fermionlike correlations, was subsequently observed in experiment both in the integrable Lieb-Liniger system (the Weiss group 2020) and previously in the corresponding lattice version, the Bose-Hubbard model (the Greiner group 2015).<sup>263</sup>

The flow of the coupling naturally leads to the concept of renormalization group (RG) flow in time  $t$ . By analogy with conventional RG ideas, increasing time plays the role of reducing the cut off with  $c = \infty$  being a strong coupling fixed point. For comparison we recall that in the usual RG picture  $c$  has scaling dimension 1 and so also flows to strong coupling. Subsequently, similar behavior was also seen in strongly coupled impurity models.<sup>264,265</sup> Extending the dynamical RG analogy one can envisage that other Hamiltonians close to the Lieb-Liniger will flow close the neighborhood of the same strong coupling fixed point, prethermalize in other words, only to end up thermalized on longer time scales if the model is not integrable, see Fig. 22. An example is provided by the lattice version of the Lieb-Liniger



**Fig. 21.** The noise correlation function  $C_2(\xi, -\xi)$ , as a function of  $\xi = x/t$  at long times for a quench from a lattice like initial state. For arbitrary values of  $c > 0$ , with  $\delta$  fixed, the system develops a distinct fermionic dip at the origin. Reprinted with permission from Iyer *et al.*, Phys. Rev. A **87**, 053628 (2013). Copyright 2013, American Physical Society.





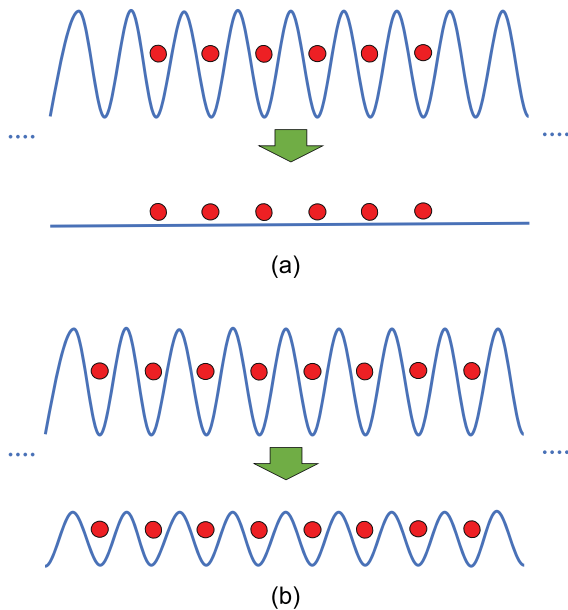
**FIG. 22.** Prethermalization in the Bose-Hubbard model. Reprinted with permission from Iyer *et al.*, Phys. Rev. A **87**, 053628 (2013). Copyright 2013, American Physical Society.

model, the nonintegrable Bose-Hubbard model which also exhibits dynamical fermionization.<sup>250</sup>

We turn now to study the global properties of the post quench system through the Loschmidt amplitude (15) and the work distribution function (16) focusing on the experimentally relevant case of a cold atom gas initially held in a deep optical lattice which is then removed entirely in the quench, see Fig. 23. The system is translationally invariant and described by the Lieb-Liniger model.

We consider  $N$  bosons on a circle of length  $L$  initially described by the state (22) with  $N$  consecutive sites filled, with  $N\delta \ll L$  so that the unfilled part of the lattice is taken to be much larger than the filled portion to avoid complications arising from the boundary conditions. Employing the Yudson resolution of the identity, the Loschmidt amplitude can be determined to be,<sup>266</sup>

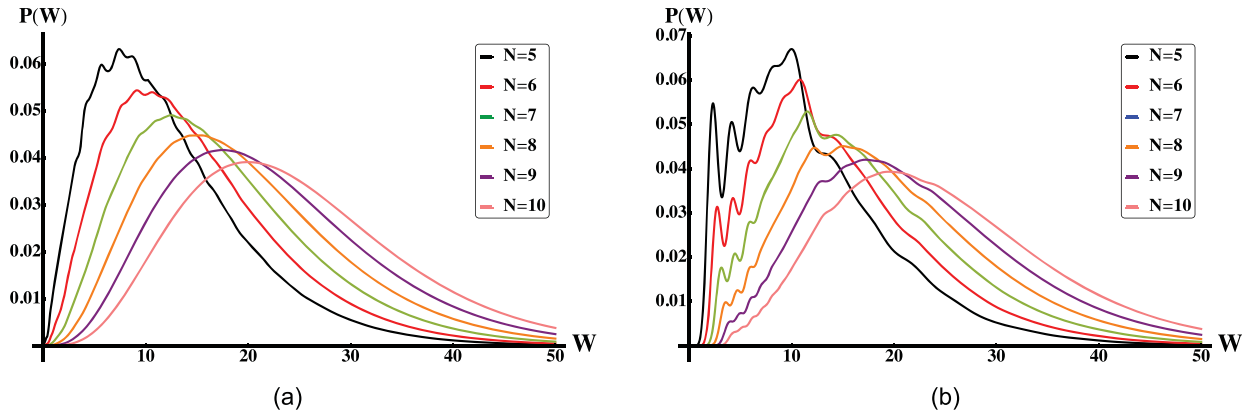
$$\mathcal{G}(t) = \left[ \frac{8\pi}{\omega} \right]^{N/2} \sum_{n_1, \dots, n_N} e^{-2/\omega [1 + i(\omega/2t)] \sum_{j=1}^N \lambda_j^2} \frac{G(\{n\})}{\mathcal{N}(\{n\})}, \quad (27)$$



**FIG. 23.** The ultracold atom gas is initially held in a deep optical lattice which is (a) completely removed—post-quench dynamics described by the Lieb-Liniger Hamiltonian or (b) merely lowered—the postquench dynamics given by Sine-Gordon Hamiltonian. Quench (b) is discussed in Ref. 269.

where  $G(\{n\}) = \det[e^{-i\lambda_j(\bar{x}_j - \bar{x}_k) - i\theta(j-k)\phi(\lambda_j - \lambda_k)}]$  and  $\theta(j-k)$  is a Heaviside function. Using the same  $1/c\delta$  expansion as before the Fourier transform of this can be explicitly found and analytic expressions for the work distribution,  $\mathcal{P}(W)$  obtained. We plot this for both noninteracting and strongly interacting bosons  $c\delta \gg 1$  in Fig. 24 for different particle number and see some commonalities as well as striking differences. Notice that the average work in both cases is the same,  $\langle W \rangle = N\omega/4$  as is the large  $W > \langle W \rangle$  behavior. The former statement can be understood from the fact that bosons are initially in non-overlapping wavefunctions and  $\langle W \rangle = \langle \Psi_0 | H | \Psi_0 \rangle$ . In comparison, the small  $W \ll \langle W \rangle$  behavior is strongly affected by the presence of interactions. Large resonant peaks are present in the interacting work distribution and can be attributed to the scattering of strongly repulsive excitations in the post quench system. Those peaks which are closest to  $\langle W \rangle$  involve fewer scattering events while those  $W=0$  involve more. As the particle number is increased these fluctuations are suppressed like  $1/\sqrt{N}$ .<sup>267,268</sup> For large systems of bosons the most interesting behavior therefore occurs in the region of  $W \sim 0$  where the effects of the interaction are most keenly felt. In this region it can be shown that the distribution decays as a power law with the exponent drastically differing between the free and interacting cases. For the former we have  $\mathcal{P}_{c=0}(W) \sim W^{N/2-1}$  whereas in the latter it is  $\mathcal{P}_{c>0}(W) \sim W^{N^2/2-1}$ , the presence of interactions in the system causing a dramatically faster decay of the work distribution. Behavior such as this will be seen in Sec. VB2 also when the excitations are gapped as well as interacting.

We can use our knowledge of  $\mathcal{P}(W)$  to investigate the global behavior of the post quench system. As a consequence of the large  $W$  agreement between the distributions for the interacting and noninteracting systems we can determine that at short times  $|\mathcal{G}(t)|^2$  is independent of the interactions. This corresponds to the initial period of expansion from the lattice in which the particles do not encounter one another. On the other hand, small  $W$  behavior provides insight to the long time dynamics, the power law decay of  $\mathcal{P}(W)$  near the origin translating to the long time power law decay of the LE. Fourier transforming the distribution for free bosons we find that as  $t \rightarrow \infty$ ,  $|\mathcal{G}(t)|^2 \rightarrow 1/t^N$  while in the interacting case we have instead  $|\mathcal{G}(t)|^2 \rightarrow 1/t^{N^2}$ , a much faster decay. We attribute this dramatic difference in the decay away from the initial state to the fact that the large repulsive interactions acting on each other forcing them to spread out into the one dimensional trap, thereby decreasing their overlap with  $|\Psi_i\rangle$ . We should note that this is true regardless of the strength of the interactions and highlights the strongly coupled nature of even weakly interacting systems in low dimensions. As we saw earlier, in the long time limit any repulsive coupling flows in time strong coupling,



**Fig. 24.** The work distribution function,  $\mathcal{P}(W)$ , for different numbers of bosons released from an optical lattice with  $\delta/m = 2$  and  $\omega = 10$ . We measure the work from  $\varepsilon_i$  the initial state energy. On top, we show the distribution for noninteracting bosons while on the bottom we show the same quantity for interacting bosons,  $c > 0$ . Reprinted with permission from Rylands and Andrei, Phys. Rev. B **100**, 064308 (2019). Copyright 2019, American Physical Society.

therefore the exponent is independent of the initial strength of  $c$ , in the TG limit ( $c = \infty$ ) one finds the same power law behavior at long times as for the finite  $c$  case. This is the dynamical fermionization discussed in Sec. V A.

The attractive regime is of significant interest. The properties of the attractive model both in and out of equilibrium are much less studied than its repulsive counterpart. This dearth of theoretical results stems from the increased complexity of the Bethe ansatz solution in the attractive model. When  $c < 0$  the model supports bound states and the ground state consists of a single bound state of all  $N$  particles.<sup>270</sup> While the eigenstates given by (18) remain valid, complex values of  $k$  which correspond to bound states are allowed. The resolutions of the identity (20) also remains formally valid provided these complex valued solutions are accounted for. A stumbling block however remains as the normalization of the Bethe states in the attractive regime is not known in closed form.

In the low density limit however it has been shown that for both repulsive and attractive interactions the spatially ordered identity (20) becomes<sup>250,255,260</sup>

$$\mathbb{1}_N = \int_{\Gamma} \frac{d^N k}{(2\pi)^N} |\{k\}\rangle \langle \{k\}|. \quad (28)$$

The contours of integration,  $\Gamma$ , lie on the real line for repulsive interactions and are spread out in the imaginary direction for the attractive case with  $\text{Im}(k_{j+1} - k_j) > |c|$ .

Making use of this here in conjunction with the same  $|c| \gg m\omega$  expansion we find that the work done in the attractive regime separates into two contributions,

$$\mathcal{P}_{c<0}(W) = \mathcal{P}_{\text{free}}(W) + \mathcal{P}_{\text{bound}}(W). \quad (29)$$

The first term  $\mathcal{P}_{\text{free}}(W)$  is the contribution from particles which do not form bound states, it is identical to the expression in repulsive case only now  $c < 0$ . The major difference imposed by this is that the effective distance between the particles is smaller  $\delta_{\text{eff}} < \delta$ , the attractive interactions promoting the clustering of particles.

The simple analytic continuation to negative coupling of the first term is reminiscent of the super Tonks-Girardeau gas. This highly

correlated state of the LL (Lieb-Liniger) model is created by preparing a repulsive LL gas in the Tonks-Girardeau limit,  $c \rightarrow \infty$  and then abruptly changing the interaction strength from the being large and positive to large and negative. The result is a metastable nonequilibrium state which exhibits enhanced correlations. Many of the properties of this state can be shown to emerge from a simple analytic continuation of the coupling to large negative values. In effect the negligible overlap of each particle of our initial state mimics the density profile of the TG gas and so super-TG like behavior is not unexpected. We should stress that the expression (28) is valid at arbitrary negative values  $c$  and so not limited to super-TG regime.

The second term  $\mathcal{P}_{\text{bound}}(W)$  is entirely different. It is due to the bound states and is calculated by deforming the contours in (28) to the real line and picking up contributions due to the poles at  $k_i - k_j = ic$  present in (18). An  $n$ -particle bound state can be shown to contribute  $\mathcal{P}_{n\text{-bound}}(W) \propto |c|^{n-1} e^{-n|c|\delta}$  with factors from multiple bound states being multiplicative.

This exponential factor means that the probability that the initial state transitions to one containing bound states is highly suppressed and in the true super-TG limit vanish entirely. Despite this, for finite  $|c|$  the bound states have a strong signature in work distribution function. Since forming a bound state will lower the energy of the system<sup>270</sup> the work distribution becomes nonvanishing at negative values of  $W$ . There is a nonzero probability that work can be extracted from the system. Importantly this does not violate the 2nd law of thermodynamics as the average work remains positive  $\langle W \rangle$ .<sup>271,272</sup> In fact, it has been observed recently that the probability of extracting work from a single electron transistor can be as high as 65% whilst still satisfying the 2nd law.<sup>273</sup>

To see this we examine the leading term of  $\mathcal{P}_{\text{bound}}(W)$  which arises due to the formation of a single two particle bound state

$$\mathcal{P}_{\text{bound}}(W) \approx N \sqrt{\frac{2\pi\omega}{m}} \frac{e^{-|c|\delta - 2W/\omega}}{\Gamma(\frac{N}{2} - 1)} \left[ \frac{2(W + \frac{|c|^2}{4m})}{\omega} \right]^{N/2-2}, \quad (30)$$

which is nonvanishing for  $-|c|^2/4m < W$ . Determining the full bound state contribution is a straightforward yet involved calculation that we will not deal with here.

## 2. The XXZ Heisenberg spin chain

The XXZ Heisenberg chain provides another example of an experimentally relevant integrable model. The Hamiltonian

$$H = J \sum_{j=1}^N \{ \sigma_j^x \sigma_{j+1}^x + \sigma_j^y \sigma_{j+1}^y + \Delta \sigma_j^z \sigma_{j+1}^z \} \quad (31)$$

models a linear array of spin interacting via anisotropic spin exchange. The isotropic case  $\Delta = 1$  is SU(2) invariant and enjoys the distinction of being the first model solved by Bethe by means of the approach that bears his name.<sup>274</sup> The generalization to the anisotropic case was given by Orbach.<sup>275</sup> The eigenstates are again characterized by a set of Bethe momenta  $\{k_j\}$  describing the motion of  $M$  down-spins in a background of  $N - M$  up-spins, and are given by

$$|\vec{k}\rangle = \sum_{\{m_j\}} \prod_{i < j} [\theta(m_i - m_j) + s(k_i, k_j) \theta(m_j - m_i)] \times \prod_j e^{ik_j m_j} \sigma_{m_j}^- |\uparrow\rangle, \quad (32)$$

where  $m_j$  the position of the  $j$ th down-spin is summed from 1 to  $N$  (the length of the chain), and the S-matrix is given by

$$s(k_i, k_j) = -\frac{1 + e^{ik_i + ik_j} - 2\Delta e^{ik_i}}{1 + e^{ik_i + ik_j} - 2\Delta e^{ik_j}}. \quad (33)$$

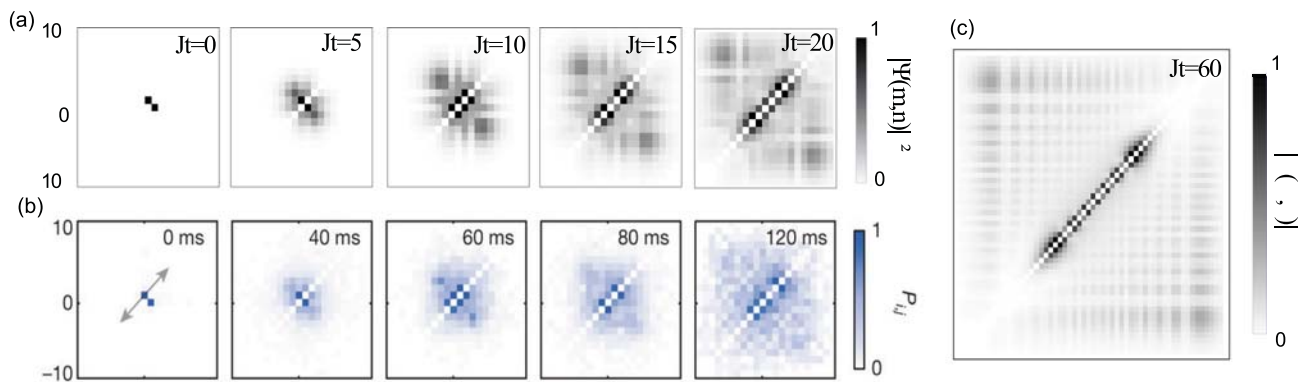
The Heisenberg chain exhibits a complex spectrum which includes bound states in all parameter regimes. To carry out the quench dynamics for the model one needs to construct the appropriate Yudson representation and use it to time evolve any initial state.<sup>276</sup> Here we display in Fig. 25 the time evolving wavefunction of two adjacent flipped spins in the background of an infinite number of unflipped spins and compare it to the experimental results (no adjustable parameters are involved.) The time evolution of the magnetization from an initial state of three flipped spins for different values of the anisotropy  $\Delta$  is given in Fig. 26. We see that excitations propagate outward after the quench forming a sharp light-cone in contrast to the Lieb-Liniger model. The boundary of the light-cone arises from the propagation of free magnons which travel with the maximum velocity

allowed by the lattice. Rays within the light-cone are the propagation of spinon bound states. As the anisotropy  $\Delta$  is increased the bound states slow down and more spectral weight is shifted to them. Due to the integrability of (31) these excitations have infinite lifetime which prevents any dispersion of these features. The introduction of integrability breaking terms can therefore be expected to alter this picture, for example through spinon decay.<sup>277</sup>

## C. Concluding remarks and outlook

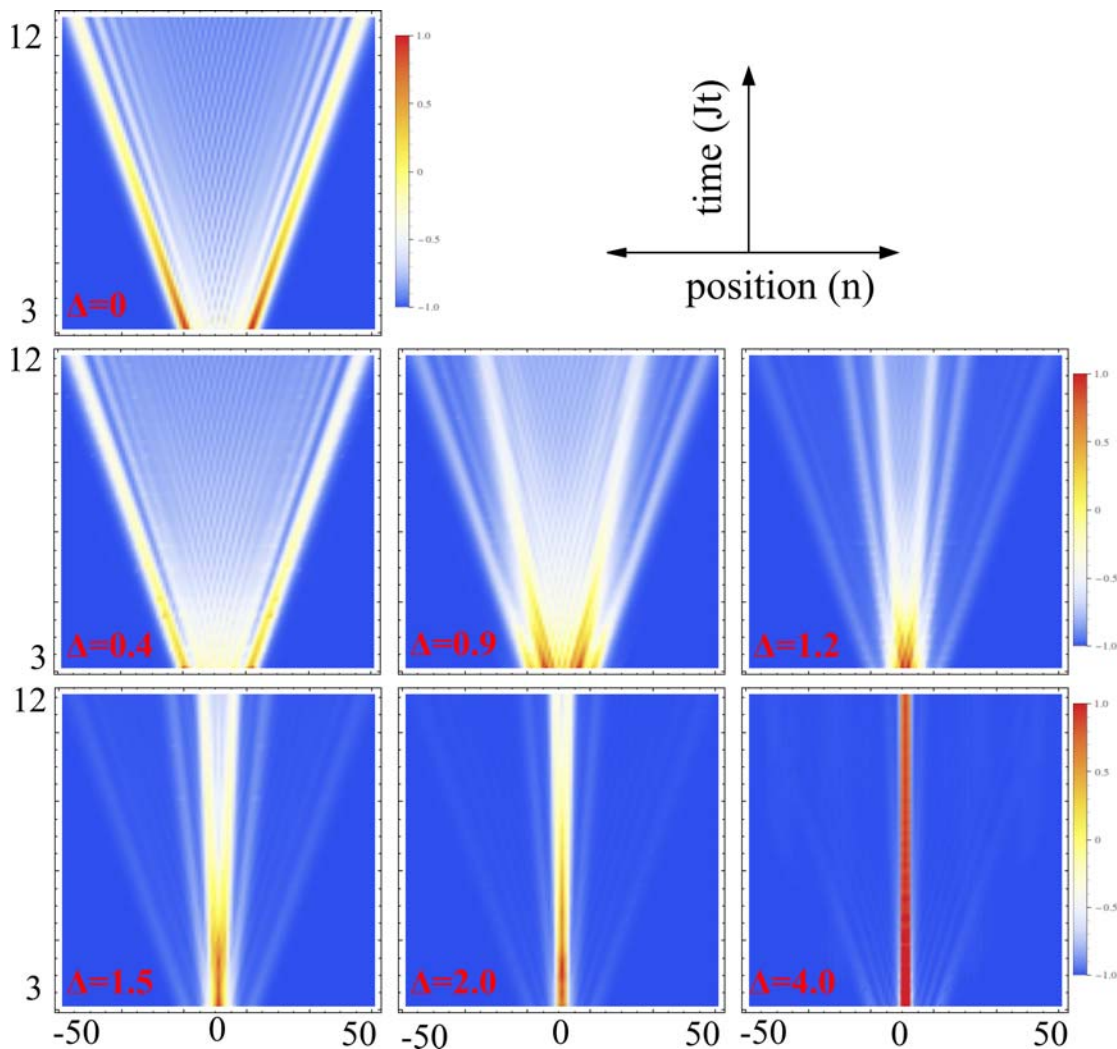
In this chapter we have explored some aspects of the far from equilibrium behavior of integrable models. After a broad overview of the current status of the field we investigated some particular phenomena through a number of illustrative examples. We saw that the Bethe ansatz solution of the Lieb-Liniger and Heisenberg models provided us with a powerful tool with which to study both the local and global, nonequilibrium behavior of these strongly coupled systems. The quench dynamics of more complex models such as the Gaudin-Yang model<sup>279,280</sup> describing multicomponent gases has also been accessed via the Yudson approach<sup>281</sup> allowing the study of phenomena such a quantum Brownian motion or the dynamics of FFLO (Fulde-Ferrell-Larkin-Ovchinnikov) states.<sup>282,283</sup> Similarly the quench dynamics of other models such as the Kondo and Anderson models are currently studied via the Yudson approach.<sup>284</sup> They give access to such quantities as the time evolution of the Kondo resonance or of the charge or heat currents in voltage or temperature driven two lead quantum dot system.

These methods we discussed could be thought as being microscopic, starting from the exact eigenstates of the system. Recently these problems have been studied from a macroscopic perspective by combining integrability and ideas from hydrodynamics.<sup>285</sup> Generalized hydrodynamics (GHD) provides a simple description of the nonequilibrium integrable models on long length scales and times. It has been utilized in studies of domain wall initial states in the Lieb-Liniger and the emergence of light-cones in quenches of the XXZ model.<sup>286,287</sup> This method allows the incorporation integrability breaking effects within the formalism, but is limited to “Euler scale” dynamics. It would be of great interest compare the results and expectations of GHD with the methods and results presented here to further



**FIG. 25.** (a) The norm of the wavefunction  $|\Psi(m, n, t)|^2$  at different times for two flipped spins initially at  $m = 1, n = 0$ . (b) The joint probabilities at different times of two spins at sites  $i$  and  $j$  initially at  $i = 1, j = 0$ , measured experimentally in Ref. 278. Reprinted with permission from Liu and Andrei, Phys. Rev. Lett. **112**, 257204 (2014). Copyright 2014, American Physical Society.





**FIG. 26.** The local magnetization after a quench from an initial state of three flipped spins at the origin for different values of the anisotropy  $\Delta$ . Time, the vertical direction, is measured in units of the exchange coupling  $J$ . Reprinted with permission from Liu and Andrei, *Phys. Rev. Lett.* **112**, 257204 (2014). Copyright 2014, American Physical Society.

understand the limitations of both the microscopic and macroscopic approaches.

## VI. NONEQUILIBRIUM PROTOCOLS FOR ONE DIMENSIONAL BOSE GASES IN ATOMTRONIC CIRCUITS

**L. Piroli, A. Trombettoni**

A promising line of research in atomtronics is the realization of configurations where several waveguides in which ultracold atoms move are merged to form circuits.<sup>5</sup> Among the challenges one has to face, an important one is the tailoring of the circuits in a way to reduce transverse instabilities during the dynamics of ultracold matter

wavepackets.<sup>28</sup> This would allow for the possibility of stable motion of the matter wavepackets across the whole circuit, including the passage through junctions and in the regions where the waveguides composing the circuit have to bend. Since transverse instabilities are suppressed in one-dimensional geometries, the lines of research of atomtronics and one-dimensional ultracold atoms have been developing tight connections in the last decade. On the one hand, the study of circuits made of one-dimensional waveguides open new directions of investigation for the community working on one-dimensional integrable systems, such as the study of junctions of one-dimensional waveguides: an example is given in,<sup>63</sup> where a junction of three Tonks-Girardeau gases is studied, and connected to the literature of coupled/intersecting nanowires. On the other hand, the amount of available results in the field of one-dimensional integrable models provides an extremely useful basis for

the characterization of ultracold matter wavepackets on such geometries, which has been at center of significant discussions in the Atomtronics@Benasque conference series.<sup>5</sup>

One-dimensional interacting bosons are well described by the integrable Lieb-Liniger model, which was extensively studied since its introduction in the sixties, also in connection with other one-dimensional integrable systems. Extensions and generalizations of the Lieb-Liniger model may apply to one-dimensional fermionic systems and mixtures, including Bose-Bose and Bose-Fermi mixtures. Therefore, the field of atomtronics circuits made of (possibly connected) one-dimensional ultracold systems is a natural arena to apply such a body of knowledge, and at the same time calls for new ideas and investigations using integrability techniques.

One-dimensional systems provide *per se* an exciting arena where, over the past decade, significant experimental technical advances have allowed for very precise studies of a series of nonequilibrium phenomena. At the same time, a number of powerful theoretical tools were developed to describe them. The study of one-dimensional systems plays a role as well in the field of atomtronics and in particular in atomtronics circuits, where matter-wave packets can be controlled and moved. When the transverse dimensions of the waveguides in which atoms move are small enough to create one-dimensional tightly confined traps and the energies involved are negligible with respect to the excitation energies of transverse degrees of freedom, then one enters the one-dimensional regime. Ultracold bosons are then effectively described by the Lieb-Liniger model,<sup>288–290</sup> belonging to the family of integrable theories. In such one-dimensional regimes quantum fluctuations play a prominent role and a general issue is whether and for what applications such one-dimensional features hamper or at variance make it easier to realize atomtronics tasks.

Here we give an account of some interesting properties of the Lieb-Liniger model and how integrability gives access to the study of some of its local and global nonequilibrium properties. The following contribution focuses on the theoretical study of two of the more useful protocols to control the quantum dynamics of the Lieb-Liniger model: (i) integrable dynamics after a quench; (ii) Floquet engineering. They are relevant for atomtronics applications, both for the possibility to have quenches and time-periodic potentials as a tool to control the dynamics and induce desired dynamical regimes and for the remarkable progress in experimental techniques enabling the possibility to vary interaction strengths, geometry of the trap and the time-dependence of the potentials acting on the atoms in one-dimensional ultracold systems.<sup>288–290</sup> In the present contribution, L.P. wrote Sec. VIA, while A.T. wrote Sec. VIB.

## A. Quench dynamics in the Lieb-Liniger model

In the early noughties, a series of cold-atomic experiments contributed to the emergence of a growing theoretical interest in the nonequilibrium dynamics of isolated quantum integrable systems.<sup>125,291</sup> For instance, in the famous “quantum Newton’s cradle” experiment,<sup>292</sup> out-of-equilibrium arrays of trapped one-dimensional (1D) Bose gases were shown not to reach thermal equilibrium within the accessible time scales. This peculiar behavior was attributed to the approximate integrability of the system: indeed, in the idealized situation where longitudinal confining potentials are neglected, a 1D gas of  $N$  bosons with mass  $m$  and pointwise interactions can be described by

the integrable Lieb-Liniger Hamiltonian.<sup>251</sup> Denoting by  $L$  the length of the system, the Hamiltonian can be written as

$$H = \int_0^L dx \left( \frac{\hbar^2}{2m} \partial_x \Psi^\dagger \partial_x \Psi + c \Psi^\dagger \Psi^\dagger \Psi \Psi \right), \quad (34)$$

where  $\Psi$ ,  $\Psi^\dagger$  are bosonic creation and annihilation operators satisfying canonical commutation relations. Here, the interaction strength is related to the one dimensional scattering length  $a_{1D}$  through  $c = -\hbar^2/ma_{1D}$ <sup>293</sup> and can be varied via Feshbach resonances<sup>294</sup> to take either positive or negative values.

Given its relative simplicity and experimental relevance, in the past decade a large number of studies have focused on the nonequilibrium dynamics in the Lieb-Liniger gas, especially within simplified protocols such as the one of a quantum quench:<sup>295,296</sup> in this setting one considers the ground state of some Hamiltonian  $H(c_0)$  ( $c_0$  denotes an internal parameter), which is suddenly changed at time  $t=0$  by an abrupt variation  $c_0 \rightarrow c$ . These studies have played an important role for the development of a general theory of integrable systems out of equilibrium.<sup>297</sup> In this section, we provide a review of a selected number of them, focusing exclusively on the simplest case of homogeneous settings (see Sec. VIC for recent further developments in the presence of confinement potentials and inhomogeneities).

## 1. The quench problem

Physical intuition suggests that after a quench an extended system should act as an infinite bath with respect to its own finite subsystems, and that local properties should relax to stationary values described by a thermal Gibbs ensemble. While for generic models this picture turns out to be correct,<sup>249,298,299</sup> a quite different scenario emerges in the presence of integrability, due to the existence of an extensive number of local conservation laws which strongly constrain the dynamics. In this case, it was proposed in Ref. 300 that the correct post-quench stationary properties are captured by a generalized Gibbs ensemble (GGE), which is written in terms of all higher local conservation laws beyond the Hamiltonian.<sup>300–302</sup> It was later discovered that quasi-local conservation laws must also be taken into account<sup>303–308</sup> and the validity of the GGE is now widely accepted.

Despite the established conceptual picture, computations based on the GGE are hard, and more generally the characterization of the post-quench dynamics remains extremely challenging in practice. In order to explain the difficulties involved, it is useful to consider the time evolution of a physically relevant observable for the 1D Bose gas, namely so-called pair correlation function<sup>309</sup>

$$g_2 = \frac{\langle \Phi | \Psi^{\dagger 2}(x) \Psi^2(x) | \Phi \rangle}{D^2}, \quad (35)$$

where  $D = N/L$  is the particle density, with  $L$  the system size, while  $|\Phi\rangle$  is the state of the system. Physically,  $g_2$  quantifies the probability that two particles occupy the same position. For a quantum quench, we have the formal expression (setting  $\hbar = 1$ )

$$\begin{aligned} \langle \Phi(t) | \Psi^{\dagger 2}(x) \Psi^2(x) | \Phi(t) \rangle &= \sum_{m,n} \langle \Phi(0) | n \rangle \langle m | \Phi(0) \rangle \\ &\times \langle n | \Psi^{\dagger 2}(x) \Psi^2(x) | m \rangle e^{-i(E_n - E_m)t}. \end{aligned} \quad (36)$$

Here we denoted by  $|n\rangle$ ,  $E_n$  the energy eigenstates and eigenvalues, respectively, while  $|\Phi(t)\rangle$  is the state of the system evolved at time  $t$  after the quench. For the Lieb-Liniger model the Bethe ansatz<sup>257</sup> is a very efficient tool to obtain most of the ingredients appearing in Eq. (36), including the matrix elements of the local operator  $\Psi^{\dagger 2}(x)\Psi^2(x)$ .<sup>310,311</sup> However, due to the complicated form of the energy eigenfunctions, there appears to be no simple way to compute the overlaps  $\langle\Phi(0)|n\rangle$  for general initial states. Furthermore, Eq. (36) involves the evaluation of a double sum over all the eigenstates of the Hamiltonian, which is currently out of reach in most of the physically interesting situations.

Due to the above difficulties, initial studies in the Lieb-Liniger model were restricted to the limit of either vanishing<sup>312,313</sup> or infinitely repulsive post-quench interactions,<sup>314–320</sup> where the Hamiltonian can be mapped onto free fermions through a Jordan-Wigner transformation. While these works already made it possible to explore in some detail interesting phenomena such as local relaxation<sup>317</sup> and “light-cone” spreading of correlation functions,<sup>312,317</sup> it remained as an open problem to provide predictions in the case of finite values of the interactions.

## 2. The quench action

A conceptual and technical breakthrough came with the introduction, by Caux and Essler, of the so-called Quench Action method,<sup>254,321</sup> which proved to be a powerful and versatile approach to the quench dynamics in integrable systems (other methods, that will not be discussed here, have also been developed, including a Yudson-representation approach, which is also suitable to study inhomogeneous initial states, see Refs. 260, 266 and the contribution of N. Andrei and C. Rylands).

It is well known that, in the thermodynamic limit, each eigenstate of an integrable system is associated with a distribution function  $\rho(\lambda)$ , where  $\lambda$  are the quasi-momenta of the (stable) quasi-particle excitations.<sup>257</sup> Based on physical arguments, it was proposed in Ref. 254 that this description could be exploited to replace the double sum in Eq. (36) with a functional integral over all distribution functions  $\rho(\lambda)$ . This approach is particularly powerful to investigate the late-time limit, for which one can write (in the thermodynamic limit)<sup>254,321</sup>

$$\lim_{t \rightarrow \infty} \langle \Phi(t) | \Psi^{\dagger 2}(x) \Psi^2(x) | \Phi(t) \rangle = \int \mathcal{D}\rho \, e^{S[\rho]} \langle \rho | \Psi^{\dagger 2}(x) \Psi^2(x) | \rho \rangle, \quad (37)$$

where  $|\rho\rangle$  denotes an eigenstate corresponding to the distribution function  $\rho(\lambda)$ . Here we introduced the “Quench Action”  $S[\rho]$ , which can be determined based on the knowledge of the overlaps  $\langle\Phi(0)|n\rangle$ . While, as we have already mentioned, it is not known how to obtain these in general, it turned out that they can be computed in several interesting cases.<sup>322–343</sup>

Given  $S[\rho]$ , the functional integral can be treated exactly by saddle-point evaluation, so that the r.h.s. of Eq. (37) can be replaced by  $\langle \rho_s | \Psi^{\dagger 2}(x) \Psi^2(x) | \rho_s \rangle$ , where  $\delta S[\rho_s]/\delta \rho = 0$ . Crucially, the saddle-point distribution function  $\rho_s(\lambda)$  determines all the post-quench local expectation values (which can be explicitly computed via exact Bethe ansatz formulas<sup>310,344–347</sup>) and thus represents an effective characterization of the late-times steady state.

The Quench Action approach was first applied in the Lieb-Liniger model for quenches from zero to positive values of the interactions,  $c_0 = 0 \rightarrow c > 0$ .<sup>348</sup> It was found that the steady state displays quantitative different features from a thermal state, unequivocally proving the absence of thermalization. The same approach also allowed for the computation of the full time evolution of  $g_2$ <sup>349</sup> (see also<sup>350,351</sup>) unveiling a quite general power-law decay to stationary values for local observables, and for a detailed study of the statistics of the work performed by the quench.<sup>352–354</sup>

## 3. Quenches to the attractive regime

In the case of quenches to repulsive interactions, the late-time steady state appears to display features that are only quantitatively different from those observed at thermal equilibrium.<sup>348</sup> In this respect, an even more interesting picture emerges for quenches to the attractive regime. These were investigated in Refs. 355 and 356 where the formalism of Refs. 254 and 348 was employed to study interaction quenches of the form  $c_0 = 0 \rightarrow c < 0$ .

The main results of these works are arguably the prediction of the dynamical formation of  $n$ -boson bound states with finite densities  $D_n$  and the characterization of the corresponding distribution of quasi-momenta  $\rho_n(\lambda)$ . Interestingly, it was shown that the value of  $n$  for which the density  $D_n$  is maximum decreases with the rescaled interaction  $\gamma = |c|/D$ . Although this result might appear counter-intuitive, there is in fact a simple physical interpretation: in the attractive regime, the bosons have a tendency to form multiparticle bound states. However, in the quench setup the total energy of the system is fixed by the initial state, while the energy of  $n$ -particle bound states increases, in absolute value, very rapidly with  $\gamma$  and  $n$ .<sup>355</sup> Therefore,  $n$ -particle bound states cannot be formed for large values of  $\gamma$ , while they become accessible as  $\gamma$  decreases.

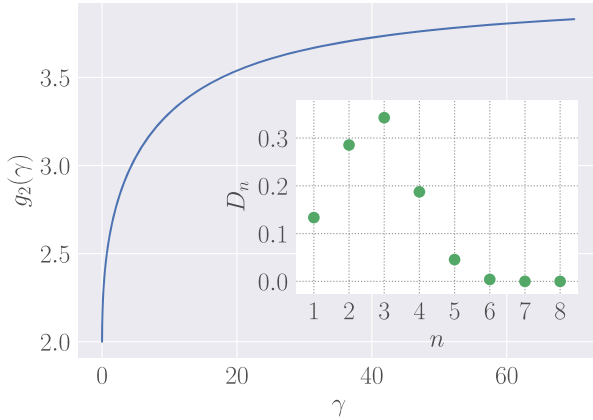
We note that the structure of the stationary state predicted in Refs. 355 and 356 is qualitatively very different from the super Tonks-Girardeau gas, which is obtained by quenching the one-dimensional Bose gas from infinitely repulsive to infinitely attractive interactions.<sup>315,357–362</sup> Indeed, the latter features no bound state, even though it is more strongly correlated than the traditional Tonks-Girardeau gas, as has been observed experimentally.<sup>359</sup> The findings of Refs. 355 and 356 are thus also interesting because they show that the physics emerging at late times after a quench depends qualitatively on the initial state of the system.

Importantly, the formation of bound states after the quench have consequences on the local correlation functions. For instance, the value of  $g_2$  at large times is always greater than 2, and increases with  $\gamma = |c|/D$ .<sup>356</sup> This is displayed in Fig. 27, and is once again qualitatively different from the case of the super Tonks-Girardeau gas. We note that these results are consistent with subsequent numerical calculations reported in Ref. 363 and based on the method developed in Refs. 364 and 365.

## B. Floquet Hamiltonian for the periodically tilted Lieb-Liniger Model

Another promising protocol for inducing and controlling interesting instances of quantum dynamics is provided by the Floquet engineering. In this scheme the original Hamiltonian—in this section the Lieb-Liniger model—is subject to a time-periodic driving  $V$ .





**FIG. 27.** Main panel: Pair correlation function, as defined in Eq. (35) in the steady state reached after a quench  $c_0 = 0 \rightarrow c < 0$ . The plot shows  $g_2$  as a function of the rescaled interaction  $\gamma = |c|/D$ , and is computed using the results of Ref. 355. Inset: Densities  $D_n$  of the  $n$ -particle bound states for the same quench, and  $\gamma = 2$ .

The Floquet Hamiltonian control then the time dynamics of the system when observed at stroboscopic times, i.e., at times multiples of the period of  $V$ . The general goal is to design  $V$  in a way that the Floquet Hamiltonian is the one inducing the desired quantum dynamics.

In general, when a periodic driving acts on an integrable model, then the resulting Floquet Hamiltonian is nonintegrable. In this section, we consider the case of the Lieb-Liniger model subject to a potential periodic in time and linear in space, which we refer to as a periodic tilting.<sup>366</sup> The Floquet Hamiltonian of the integrable Lieb-Liniger model for such linear potential with a periodic time-dependent strength is integrable and its quasi-energies can be determined using well known results for the undriven Lieb-Liniger model.

We pause here to comment about the relevance of the investigation of Floquet engineering, and periodic tilting in particular, starting from the Lieb-Liniger Hamiltonian for atomtronics applications and perspectives. Controlling matter-wave dynamics in waveguides and other atomtronics circuitry and components is in general an interesting perspective to be discussed and studied. A time-independent potential linear in space induces a motion in the atomtronics devices, and a time-dependent periodic tilt can be used to control the motion across, to and fro, a circuit. As discussed in the Introduction, to reduce transverse excitations it may be convenient to use and merge one-dimensional waveguides, and a natural question is what is the effect of a time-dependent periodic tilting in such one-dimensional systems.

We then consider the periodic tilting

$$V(x, t) = f(t) x$$

with  $f(t)$  a periodic function with period  $T$ . The Lagrangian density of the system is

$$\mathcal{L} = \frac{i}{2} (\Psi^\dagger \partial_t \Psi - h.c.) - \frac{1}{2m} \partial_x \Psi^\dagger \partial_x \Psi - \frac{c}{2} \Psi^\dagger \Psi^\dagger \Psi \Psi - V(x, t) \Psi^\dagger \Psi, \quad (38)$$

where  $h.c.$  denotes the hermitian conjugate of the first term and  $\Psi = \Psi(x, t)$ .

When the potential  $V$  is time-independent with  $f(t)$  constant, then it is well known that one can gauge away the potential linear in space by moving to the center-of-mass accelerating frame. Notice that this property is valid in any dimension and also for interacting systems, as long as the two-body interaction depends only on the relative distance (for a pedagogical presentation see, e.g., Ref. 367).

Let now come back to the case of  $f(t)$  periodic in time. Proceeding as one does for the single-particle and the two-particles cases,<sup>366,368</sup> one can solve the Schrödinger equation of the many-body interacting model. To this aim, one introduces the following gauge transformation:

$$\Psi(x, t) \equiv e^{i\theta(x, t)} \varphi(y(t), t), \quad (39)$$

where

$$y(t) = x - \xi(t),$$

with the functions  $\xi(t)$  and  $\theta(x, t)$  to be suitably determined in order to gauge away the potential term  $V$  from the Lagrangian density when rewritten in terms of the field  $\varphi$ .

The functions  $\xi$  and  $\theta$  are determined as it follows. We start by imposing

$$\partial_t \xi = \frac{1}{m} \partial_x \theta \quad (40)$$

and

$$-\partial_t \theta = \frac{1}{2m} (\partial_x \theta)^2 + x f(t). \quad (41)$$

We now make the ansatz

$$\theta(x, t) = m x \partial_t \xi + \Gamma(t), \quad (42)$$

finding the conditions

$$m \partial_t^2 \xi = -f(t) \quad (43)$$

and

$$\partial_t \Gamma = -\frac{m}{2} (\partial_t \xi)^2, \quad (44)$$

determining  $\xi(t)$  and  $\Gamma(t)$  in terms of  $f(t)$ . From the differential equations (43)–(44) one gets<sup>366</sup>

$$\theta(x, t) = -x \int_0^t f(\tau) d\tau - \frac{1}{2m} \int_0^t \left[ \int_0^\tau f(\tau') d\tau' \right]^2 d\tau. \quad (45)$$

Notice that, with our choices of the initial conditions [ $\xi(0) = d\xi(0)/dt = 0$  and  $\Gamma(0) = 0$ ], one has  $\theta(x, 0) = 0$  and  $y(0) = x$ . Using (45), from (40),  $\xi$  can be readily determined.

For the sake of simplicity we will discuss the case

$$\int_0^T f(\tau) d\tau = 0 \quad (46)$$

(referring to<sup>366</sup> for a discussion about the case  $\int_0^T f(\tau) d\tau \neq 0$ ). The major simplification is that the gauge phase (45) does not depend anymore, at stroboscopic times, on the spatial variable, i.e.,  $\theta(x, T) \equiv \theta(T)$ .

Provided the condition (46) holds, with the functions  $\theta$  and  $\xi$  previously determined we can rewrite the Lagrangian density (38) in terms of  $\varphi(y, t)$  which involves no longer the external potential:

$$\mathcal{L} = \frac{i}{2} (\varphi^\dagger \partial_t \varphi - h.c.) - \frac{1}{2m} \partial_y \varphi^\dagger \partial_y \varphi - \frac{c}{2} \varphi^\dagger \varphi^\dagger \varphi \varphi. \quad (47)$$

Notice that the outlined procedure also works for a more general potential of the form  $V(x, t) = xf(t) + g(t)$ .

To determine the Floquet Hamiltonian we need to determine the function  $\theta$  at the stroboscopic times:  $t \equiv nT$ , with  $n$  integer. One has

$$\theta(nT) = -\frac{1}{2m} \int_0^{nT} dt \left[ \int_0^t dt' f(t') \right]^2 \quad (48)$$

and

$$\xi(nT) = -\frac{1}{m} \int_0^{nT} dt \int_0^t dt' f(t'). \quad (49)$$

Now we want show that the ratios  $\frac{\xi(nT)}{nT}$  and  $\frac{\theta(nT)}{nT}$  do not depend on time, i.e., on  $n$ . Let be  $\mathcal{F}(t)$  a function such that  $\frac{d\mathcal{F}}{dt} = f(t)$ . The constant of integration is chosen to be such that  $\mathcal{F}(0) = 0$ . From (46) one has

$$\int_0^T f(t) dt = \mathcal{F}(T) = 0, \quad (50)$$

so that one can see that  $\mathcal{F}(t)$  is a periodic function of period  $T$ . Using the definition of the function  $\mathcal{F}$ , from (49) one gets

$$\xi(nT) = -\frac{1}{m} \int_0^{nT} \mathcal{F}(t) dt = -\frac{I}{m} n, \quad (51)$$

where  $I \equiv \int_0^T \mathcal{F}(t) dt$ . It follows that the ratio  $\frac{\xi(nT)}{nT}$  is  $n$ -independent. The same reasoning applies for the gauge phase  $\theta$ , since it is

$$\theta(nT) = -\frac{I'}{2m} n, \quad (52)$$

where  $I' \equiv \int_0^T \mathcal{F}^2(t) dt$ .

We are now able to write the Floquet Hamiltonian, which is found to be

$$H_F = \sum_{j=1}^N \left( \frac{\hat{p}_j^2}{2m} + \frac{\xi(T)}{T} \hat{p}_j - \frac{\theta(T)}{T} \right) + c \sum_{j<i} \delta(x_j - x_i). \quad (53)$$

We observe that the previous derivation of the Floquet Hamiltonian is valid not only for the one-dimensional Lieb-Liniger Hamiltonian, but also for a generic interacting system in any dimension subject to a periodic tilting. In the one-dimensional Lieb-Liniger this result implies the main point of this section, relevant for atomtronics application: the Floquet Hamiltonian (53) is *integrable*, as it can be immediately seen. A further important comment, on which we are going to comment more in the following, is that the derivation of (53) is valid for translational invariant systems.

One can then apply the standard Bethe ansatz techniques (see<sup>251,257</sup>) to compute the quasi-energies and the eigenfunctions. More precisely, one has to compute the pseudo-momenta  $k_j$  obeying the Bethe equations. If the system is subjected to periodic boundary conditions, the pseudo-momenta  $k_j$  are determined in terms of the following Bethe equations:

$$k_j L + 2 \sum_{i=1}^N \arctan\left(\frac{k_j - k_i}{mc}\right) = 2\pi \left(j - \frac{N+1}{2}\right), \quad (54)$$

for  $j = 1, \dots, N$ , where  $L$  is the circumference of the ring in which the system is confined.

Using the previous results one can write the many-body states at the stroboscopic times. A multiparticle  $|\psi\rangle$  state for the Lieb-Liniger model read (apart from the normalization factor)

$$|\psi\rangle = \int d^N x \chi(x_1, \dots, x_N, t) \Psi^\dagger(x_1, t) \cdots \Psi^\dagger(x_N, t) |0\rangle \quad (55)$$

where  $\chi(x_1, \dots, x_N, t)$  is the  $N$ -body wavefunction.

The wavefunction  $\chi$  in (55) is a solution of the Schrödinger equation

$$i\partial_t \chi(x_1, \dots, x_N, t) = H \chi(x_1, \dots, x_N, t), \quad (56)$$

with  $H$  being the Lieb-Liniger Hamiltonian in first quantization

$$H = -\frac{1}{2m} \sum_{j=1}^N \partial_x^2 + c \sum_{j<i} \delta(x_j - x_i) + \sum_{j=1}^N V(x_j, t). \quad (57)$$

Using (39) one can write for the periodically driven model

$$|\psi\rangle = \int d^N y \eta(y_1, \dots, y_N, t) \varphi^\dagger(y_1, t) \cdots \varphi^\dagger(y_N, t) |0\rangle. \quad (58)$$

The relation between the functions  $\chi$  and  $\eta$  is given by

$$\chi(x_1, \dots, x_N, t) \equiv \prod_{i=1}^N e^{i\theta(x_i, t)} \eta(y_1, \dots, y_N, t), \quad (59)$$

where  $\eta$  is the solution of the same Schrödinger equation (56) but with  $n$  external potential ( $V = 0$ ).

An important comment is related to the fact that, as mentioned, the treatment of the Lieb-Liniger model in a periodic tilting presented in this section is valid only for translational invariant systems. However, for setups relevant for atomtronics one has to consider the effect of their particular boundary conditions. As an example, one can consider circuits obtained merging one-dimensional waveguides. The derivation presented here does not longer applies and the results of this section provides only a first step toward the determination of the correct Floquet Hamiltonian, a study which in the opinion of the author is a deserving subject in view of the possible obvious applications in atomtronics components and circuits. Separate considerations apply to ring geometries. One can think to periodically rotate the ring, with now the angle  $\varphi$  playing the role of the coordinate  $x$  since one can show<sup>366</sup> that *in the comoving reference frame* (and under the assumption that  $f(t)=0$  at the stroboscopic times) the Floquet Hamiltonian has the form (53). This is analogous of what occurs for a two-dimensional harmonic potential in rotation, where the Hamiltonian in the rotating frame has the form  $H - \Omega L_z$ <sup>121</sup> and one can slightly deform the isotropic potential to break translational invariance. The equivalent of this in a periodically rotating ring geometry could be the addition of an out-of-plane periodic rotation component.

### C. Concluding remarks and outlook

The past few years have witnessed very rapid developments within the theory of integrable systems out of equilibrium. Arguably, the most important piece of progress pertains the introduction of the so-called generalized hydrodynamics (GHD).<sup>285,369</sup> This is a very

powerful framework, which builds upon the techniques developed in the idealized case of homogeneous systems, and allows one to provide exact predictions also for inhomogeneous settings, although only at hydrodynamic scales.

While a review of these results is beyond the scope of the present article, we note that recent works have shown that GHD is more than adequate to tackle exactly experimentally relevant setups of repulsive 1D Bose gases, including systems with confining potential,<sup>370–373</sup> spatial inhomogeneities<sup>374</sup> and dephasing noise.<sup>375</sup> In fact, quite remarkably, GHD predictions have now also been experimentally verified by monitoring clouds of bosonic cold atoms trapped on an atom chip.<sup>376</sup>

It would be extremely interesting to extend some of these recent results to inhomogeneous 1D Bose gases with attractive interactions, where the study of homogeneous quantum quenches have already revealed unexpected new features. More generally, a promising route is to analyze the out-of-equilibrium dynamics of even more complicated inhomogeneous integrable quantum gases, such as multicomponent mixtures of fermions and bosons,<sup>377,378</sup> for which the emergence of interesting phenomena at the hydrodynamic scale has been already pointed out in simple settings.<sup>380,383</sup>

Going beyond quench protocols, the effect of a time-periodic tilting in the Lieb–Liniger model with repulsive interactions has been discussed. It was shown that the corresponding Floquet Hamiltonian is integrable, by studying the spectrum of the quasi-energies and the dynamics of the system at stroboscopic times. Importantly, the analysis presented for the Lieb–Liniger model can be extended to other 1D integrable systems in time-periodic linear potentials such as, for instance, the Yang–Gaudin model for fermions. In the future, it would be very interesting to study the effect of periodic tilting in more general configurations. A main issue to be studied starting from the results presented here is that of the boundary conditions of the specific atomtronics system of interest when subject to a periodic tilting. Among the different cases that would provide a worthwhile investigation is that of atomtronics circuits periodically tilted and their application to atomtronics tasks.

## ACKNOWLEDGMENTS

We are grateful to P. Calabrese, A. Colcelli, F. Essler, G. Mussardo, and G. Sierra for several enlightening discussions on the subjects presented here and on related topics. Further acknowledgments go to the participants to the first week of the 2019 Atomtronics conference in Benasque, and in particular to V. Ahufinger, V. Bastidas, D. Cassettari, J. Mompart, N. Proukakis, F. Scazza, and W. von Klitzing.

## VII. PERSISTENT CURRENTS AND VORTICES IN ATOMTRONIC CIRCUITS

**T. Bland, M. Edwards, N. P. Proukakis, A. Yakimenko**

Atomtronics relies on the flow of coherent matter waves in the form of atomic Bose–Einstein Condensates (BECs) in closed circuits, such as in the form of closed toroidal traps, or more extended, race-tracklike, potentials. Persistent currents in such geometries enable fundamental studies of superfluidity and may lead to

applications in high-precision metrology and atomtronics.<sup>5,145</sup> The question of the generation and stability of the atomic persistent currents – which in the absence of external driving should be topologically protected – is of fundamental importance; thus it has been the subject of numerous experimental and theoretical investigations<sup>12,16,41,67,93,140,142,143,161,381–394</sup>

The quantized circulation in a ring effectively corresponds to an  $m$ -charged vortex line pinned at the center of the ring-shaped condensate, where the vortex energy has a local minimum. Noting that there is no condensate density at that location, we can think of this as a “ghost” vortex – in the sense that, at some radial distance from the center of the closed loop where there is non-negligible superfluid density, the arising phase profile is identical to that corresponding to a vortex located at the center, where the superfluid density is practically zero. Since the vortex line energy increases with condensate density such a vortex turns out to be bounded by the potential barrier, that is why even multicharged ( $m > 1$ ) metastable vortex states can be very robust. The generation and decay of a persistent current is governed by the dynamics of these quantum vortices, which can be deterministic, or random, depending on the particular setting considered.

Specifically, as modeled theoretically and observed experimentally, persistent currents can form in toroidal BECs by stirring the condensate with an optical paddle potential, imparting angular momentum in the ring through the generation of vortices and through the decay dynamics after an external perturbation.<sup>12,16,41,67,161,382,383,386–388,390,391,395</sup> They can also be induced by transmitting angular momentum from a Laguerre–Gaussian beam.<sup>140,142,389</sup> Moreover, persistent currents can also spontaneously form in toroidal BECs as phase defects appearing after a quench into the BEC state.<sup>93,392–394</sup> Persistent currents also arise in multicomponent condensates in a ring geometry.<sup>143,385,396–398</sup>

Coupled persistent currents of ultracold atomic gases provide a possibility to investigate the interaction of the superflows in a tunable and controllable environment, providing the possibility for precision measurements and even potentially controllable quantum gate operations. Previous theoretical studies<sup>399–401</sup> have drawn considerable interest to systems of coupled circular BECs. Using accessible experimental techniques, it is possible to consider a variety of physical phenomena in this setting: from Josephson effects in the regime of weak interactions (where the superflow decays by inducing phase slips reviewed in Sec. VIII) to quantum Kelvin–Helmholtz instability for merging rings.

In this contribution we review recent developments in the understanding of the formation and dynamics of persistent currents in such closed geometries based on mean-field, dissipative, and stochastic simulations. We start by considering the mechanism of formation of persistent currents in a racetrack BEC, induced by a stirring potential (Sec. VII A) (which also encompasses ring-trap geometries as a special case). We then discuss more complicated atomtronic architectures, focusing on BECs trapped in two coupled toroidal potentials which are either embedded within a single plane, or are linked transversally. Specifically, we firstly review (Sec. VII B) recent work<sup>394</sup> discussing the spontaneous formation of persistent currents in coplanar double ring geometries. We then present a brief overview (Sec. VII C) of recent investigations<sup>402–404</sup> of the dynamics of quantum vortices in a pair of vertically stacked atomtronic circuits. We end with some concluding statements.



## A. Mechanism for producing flow in a racetrack atom-circuit BEC by stirring

### 1. How stirring a racetrack atom circuit produces flow

We start by presenting a picture of how macroscopic flow is produced in a BEC confined in a racetrack atom-circuit by stirring with a wide rectangular barrier within the Gross-Pitaevskii model. The atom-circuit BEC is strongly confined to a horizontal plane and the 2D racetrack channel potential (see Fig. 28) consists of two half-circles separated by straightaways of length  $L$ ; we note that in the limit  $L = 0$ , this reduces to a ring potential, so the present discussion fully encompasses that setting.<sup>405</sup>

It is well-established that flow is accompanied by production and motion of vortices<sup>67,161,382,383</sup> or dark solitons<sup>406</sup> in the condensate. Here, we describe how and where vortices form, how they move thereafter, how circulation localized to a vortex is converted into macroscopic flow around the entire racetrack, and what conditions lead to the final amount of flow.

### 2. Creation of a single unit of flow: Vortex swap

Flow can be induced in a racetrack BEC atom circuit by stirring with a weak-link potential barrier. As the stirring barrier moves and strengthens, it produces a region of lowered density. This region of depressed density causes a backflow (flow opposite the stir direction) to develop in this region. This backflow spawns a vortex (circulation same as the stir) located on the outside of the channel and an antivortex (antistir circulation) on the inside. At a critical value of the barrier height, the two vortices swap positions. This event generates two disturbances that move away from the barrier in opposite directions at the average speed of sound. The first is the vortex/antivortex pair that moves off in the antistir direction and the second is a compression wave moving in the stir direction.

This backflow is illustrated in Fig. 29(a). In the full figure, we have plotted a series of snapshots of the velocity distribution from shortly before until shortly after the creation of the first unit of flow. It is easy to see that the backflow speed is greatest at the inner and outer

edges of the channel where the racetrack plus barrier potential is largest. As the barrier moves, the condensate in front of the barrier must migrate to the back of the barrier. The atoms at the inner and outer channel edges must move faster to avoid the regions of high potential. In this way vortices, are formed by stirring with a barrier that is much wider than the stirred condensate.

When the height of the barrier reaches a critical value, the vortex migrates from the outside to the inside of the channel as can be seen in Fig. 29(b). Shortly after this vortex swap two disturbances are generated. The first is the vortex/antivortex pair, located on the inside and outside of the channel, respectively, move away from the barrier in the antistir direction. This vortex pair causes atoms on the antistir side of the barrier to flow in the stir direction between the vortices. The second disturbance is a compression wave that propagates away from barrier region in the stir direction. This compression wave also moves atoms in the stir direction. Both disturbances move at a speed that is approximately the local speed of sound [ $c(\mathbf{r}) = \sqrt{gn_c(\mathbf{r})/m}$ ] averaged over the cross section of the condensate. These disturbances are the mechanism by which the localized circulation in the form of a vortex is converted into macroscopic flow around the entire racetrack.

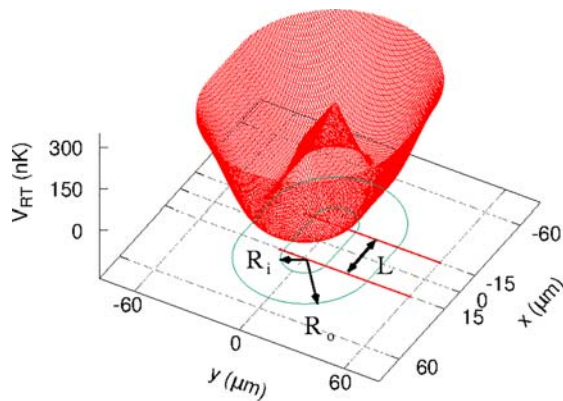
### 3. Final flow production: Flow overtakes the barrier

The final circulation produced can be roughly predicted as the number of units of quantized flow that lies closest to the speed of the stirring barrier. The exact number depends on the details of the stirring and the geometry of the racetrack as we describe below. When vortices inside the racetrack potential are generated the circulation they provide is localized near their cores. As stirring proceeds, this circulation is converted into a nearly constant tangential velocity component around the midline track by the pair of disturbances generated each time a vortex swap occurs.

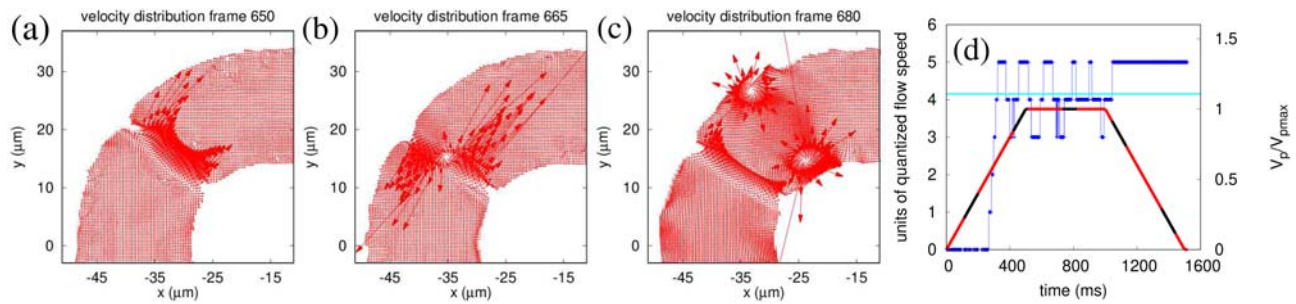
The circulation as a function of time is shown in Fig. 29(d) for the case where  $L = 30 \mu\text{m}$ ,  $v_{\text{stir}} = 339 \mu\text{m/s}$  and,  $T = 0 \text{ nK}$ . This graph shows that the circulation (shown as the blue curve) is zero until a succession of vortex-swap events produces enough flow so that the flow generated is greater than the speed of the stirring barrier (shown as the cyan horizontal line in the figure in quantized flow speed units).

In this case, five units of flow exceeds the barrier speed by almost a full flow speed unit. When the disturbance pair generated by the first vortex swap travels around the racetrack and comes back to sweep through the barrier region again, they cause a *forward flow* to develop in the barrier region. At this moment, an inverse vortex swap event can occur causing the total circulation to decrease by one unit. In this way, the circulation can oscillate around the number of units that makes the flow closest to the stir speed of the barrier.

Another circulation-changing mechanism that is only present in the non-ring racetrack case occurs when the barrier transitions from straight parts of the racetrack to curved parts or vice-versa. The times when the barrier is on straight or curved parts are indicated in Fig. 29(d) by the red- and black-colored curve that depicts the barrier height. This graph is colored red for times when the barrier is on the curved parts of the racetrack and black-colored when it is on the straightaways. Careful examination of the circulation graph shows that, when the barrier transitions from curved to straight (red to black) racetrack parts, the circulation increases by one unit. When the barrier transitions from straight to curved parts the circulation decreases by



**FIG. 28.** The atom-circuit racetrack potential consists of two semicircular endcaps (inner radius  $R_i = 12 \mu\text{m}$ , outer radius  $R_o = 36 \mu\text{m}$ ) separated by straightaways of length  $L$ . This figure was created using data reproduced from Eller *et al.*, Phys. Rev. A **102**, 063324 (2020). Copyright 2020, American Physical Society.



**FIG. 29.** Velocity distributions of the racetrack condensate during the ramp-up of the stirring. (a) A backflow plus vortex/antivortex pair develops in the barrier region, (b) the vortex/antivortex pair swap, (c) the vortex/antivortex pair move away from the barrier in the antistir direction while a compression wave moves off in the stir direction. (d) Circulation (blue line) around the midline track versus time for this case. The red and black curve gives the barrier height versus time. The curve is colored red when the barrier is on the curved parts of the racetrack and colored black when it is on the straight parts. The quantity  $V_{p,max}$  is the maximum energy height of the barrier during the stir. The straight cyan curve shows the stirring speed of the barrier in units of the quantized flow speed,  $v_{flow} = 2\pi\hbar/(Ms)$ , where  $s$  is the arc length of the racetrack midline. Parameters:  $L = 30 \mu\text{m}$ ,  $v_{stir} = 339 \mu\text{m/s}$ . Results shown are based on zero-temperature Gross-Pitaevskii mean-field theory. Panel (d) of this figure was created using data reproduced from Ref. 417.

one unit. We also note that this only happens when the barrier strength is above a critical value.

The general mechanism for flow production in the racetrack by stirring with a rectangular barrier in the context of the Gross-Pitaevskii equation can thus be summarized as follows. The stirring barrier both moves and increases in strength. This generates a backflow in the region of depressed density. This backflow is fastest at the inner and outer channel edges in this region. This flow spawns a vortex/antivortex pair at the outer and inner edges, respectively. Eventually these two vortices swap locations generating a moving vortex (now on the inner channel edge)/antivortex (now at the outer edge) pair that moves away from the barrier in the antistir and also generating a compression wave that moves away from the barrier in the stir direction. These disturbances both move at the average speed of sound. The total amount of flow produced is roughly the number of flow-speed units closest to the speed of sound.

## B. Persistent currents in coplanar double-ring geometries

Having identified the key mechanism for flow production in the context of a pure  $T=0$  BEC, it is natural to also consider the role of phase fluctuations and dissipation on the (spontaneous) emergence of supercurrents, and what happens when multiple ring-trap geometries are coupled.

### 1. Spontaneous persistent current formation in a ring trap

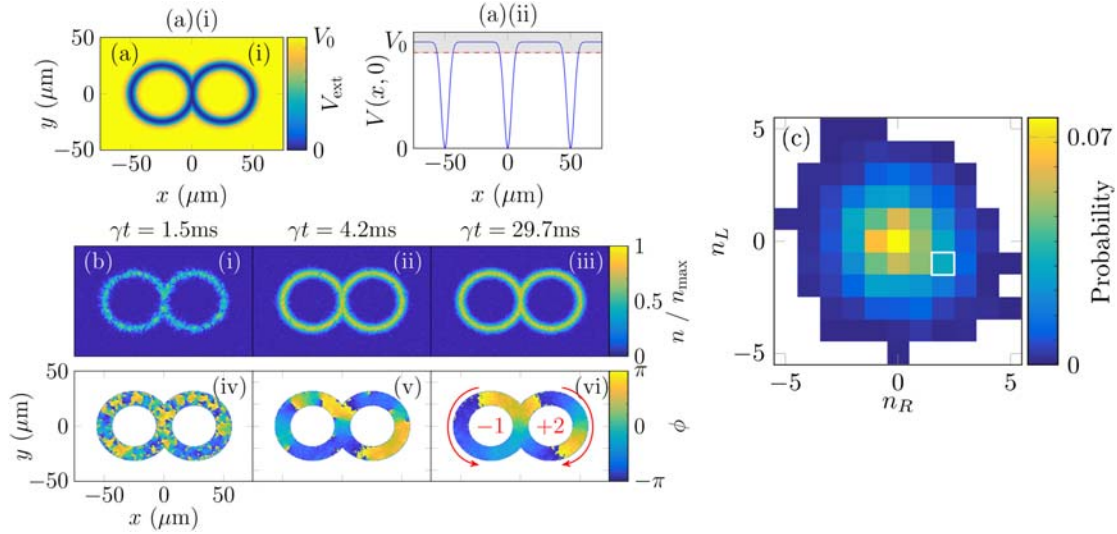
The formation of persistent currents in a ring trap can also proceed spontaneously; it is well-known that the generation of a superfluid in such a geometry can carry with it a randomly generated winding number, which is expected to be statistically distributed about the most probable “zero” value (which corresponds to the absence of a persistent current).<sup>392,393</sup> This is because phase coherence forms locally in a ring, and the size and width of the toroidal geometry, along with the rate of the actual quench leading to the formation of the ring-trap condensate, control the maximum winding number that can spontaneously emerge.<sup>386,394</sup>

This is already well-known in the context of the Kibble–Zurek mechanism,<sup>407,408</sup> which relates the generated winding number to the quenching rate of the driven phase transition, an effect already discussed and observed across different physical systems. Among those, this effect has also been confirmed in ultracold atoms in ring-trap geometries through a controlled gradual cooling rate quench, producing an experimentally observed distribution of winding numbers,<sup>392</sup> in agreement with numerical and theoretical expectations.<sup>394</sup> In fact, the local nature of such coherence evolution implies that this effect of spontaneous persistent current generation already manifests itself even in the limit of very rapid (or instantaneous) quenches toward a coherent superfluid regime. The existence of phase fluctuations and nonzero winding numbers can, for example, affect the dynamics of otherwise deterministically-generated dark solitons in ring-trap geometries.<sup>409</sup>

### 2. Spontaneous persistent current formation in coplanar connected ring traps

Next, we turn our attention to the dynamics of winding numbers in connected geometries, focusing here on the simplest possible such coplanar example, based on the 2D geometry shown in Fig. 30(a). Utilizing state-of-the-art numerical simulations based on the stochastic (projected) Gross-Pitaevskii equation,<sup>393,394,410–417</sup> we find that—as in the case of a single ring trap—coherence within the double-ring trap forms locally during condensate growth, as shown in Fig. 30(b).

After a quench, the phase develops locally along and across the ring circumference,<sup>393</sup> with an early such example of typical evolution shown by the phase profile of Fig. 30(b-iv). Although at the common interface around  $x \sim 0$  (where the trap depth has the same depth and width as the outer double-ring edges) the phase of the superfluid is constrained to be the same in both connected rings, this does not nonetheless dictate the behavior of the emerging phase in the remaining “unconnected” regions forming the bulk of the ring’s spatial extent. Specifically, the unconnected regions in the double-ring geometry are free to randomly establish their own phase dynamics (constrained by the size and width of the unconnected regions), thus often leading to nonzero winding numbers with varying (nonlinear) phase gradients across the ring circumference.



**FIG. 30.** (a) Simplest coplanar connected double-ring geometry: Shown are (i) the 2D potential, and (ii) the cut through  $y = 0$ . Note that the interfacial potential considered here is identical to that of the outer unconnected regions of the rings. (b) Formation dynamics of characteristic numerical realization with different winding numbers across the two rings: shown are growing density (from a noisy initial configuration; top row (i)–(iii)) and corresponding phase evolution [bottom row, (iv)–(vi)], for a characteristic example with distinct winding numbers  $n_{\text{left}} = -1$  and  $n_{\text{right}} = +2$  across the left and right rings, respectively. (c) Histogram of all possible steady-state winding number contributions, when performing numerical quenches from an initial noisy configuration. The indicated white box highlights the case  $(n_{\text{left}}, n_{\text{right}}) = (-1, +2)$  considered in (b). We have explicitly verified that once such an unequal winding number contribution forms after sufficient system relaxation/equilibration, it does remain stable for all subsequent evolution (with this feature also found to be insensitive to the precise choice of the “growth” parameter  $\gamma$  in the stochastic simulations). Parameters:  $N \sim 2 \times 10^5$   $^{87}\text{Rb}$  atoms,  $T = 10$  nK, effective 2D interaction strength  $\tilde{g} = 0.077$ ; Tight harmonic transverse confinement with  $\omega_z = 2\pi \times 1000$  Hz, with the 2D potential defined by  $V(x, y) = V_0 \min(1 - \exp[-2(\rho(x - R, y) - R)^2/w^2], 1 - \exp[-2(\rho(x + R, y) - R)^2/w^2])$ , where  $\rho(x, y) = \sqrt{x^2 + y^2}$ ,  $V_0 = 1.1\mu$ , the ring radius is  $R = 25\mu\text{m}$  and its width is  $w = 6\mu\text{m}$ . Adapted with permission from Bland *et al.*, J. Phys. B. **53**, 115301 (2020). Copyright 2020, Author(s) licensed under a Creative Commons Attribution 4.0 License.<sup>394</sup>

An example of such a long-term behavior with winding numbers  $-1$  and  $+2$  across the left and right rings, respectively, is shown in Fig. 30(b–vi) with a positive winding number referring arbitrary, by convention,<sup>387,394</sup> to clockwise rotation. In the double-ring case, we also find a distribution of winding numbers about the most probable value of zero net winding number, as shown in Fig. 30(c). In fact, when integrating over the winding numbers of the other ring, we find that (in our chosen, experimentally-relevant, geometry), the distribution of winding numbers in each ring actually exactly matches that of the single ring trap with the same radius, width and depth.<sup>394</sup> We expect this to be true for the majority of experimentally-relevant potentials, for which the ring radius typically largely exceeds any transversal width (and motion is frozen out in the third, transverse, direction).

Remarkably, our previous work<sup>394</sup> has shown such features to be largely independent of the exact details of the connected geometry, provided it does consist of two (2D) planar-connected closed geometries with a unique single (extended) connected region. For example, we have verified all above conclusions to be also valid in a figure-eight (“lemniscate”) potential, where there is a real crossing in the path of propagating ultracold atoms.<sup>394</sup> By extension, we would, therefore, expect similar features to hold in extended or “flattened” geometries, such as connected race-track geometries, as the underlying physics is that of how much winding can be supported by the combination of loop radius  $R$  and width  $w$ , which are found to obey the winding number relation  $\langle |n_w| \rangle \sim \sqrt{2\pi R/w}$ .<sup>393,394,408</sup>

Given the potential independence of winding numbers supported across two identical connected ring traps, it is interesting to enquire

about the stability of such features. Essentially, one can think of a winding number of, say,  $\pm n$  (where  $n = 0, 1, 2, \dots$ ) around a closed loop (whether exactly ring-shaped, or not), as being due to the existence of a “ghost” vortex trapped in the middle of the closed loop. Using such an intuitive interpretation, the winding number of a ring trap will change by an integer unit if such a “ghost” vortex is allowed to leave its enclosure, mapped out by the underlying trap potential. Due to the topological protection of the winding number, such an effect can be achieved by deforming the system topology through a change in the trap potential: for example, in the single ring-trap case, this could be achieved by opening a small hole in the potential, such that the enclosed ghost vortex can escape to the region outside of the ring. The related topic of phase slips in the presence of fluctuations is discussed in Sec. VIII.

In the double-ring geometry, we have explicitly verified that the transfer of the winding number from one side of the double-ring geometry, to the other can be facilitated by allowing for a zero-potential region to connect the two sides. Such a transfer can be reasonably controlled by the specific details of the potential deformation, even potentially leading to the annihilation of oppositely-oriented superflows (corresponding to ghost vortices of opposite circulation, which can hence annihilate), a topic of active ongoing research investigations to harness potential atomtronic applications. Our present work for interacting superfluids adds to that of tunneling angular momentum states considered at the single-particle level in single-component condensates<sup>418–420</sup> and also for two-component condensates.<sup>396</sup>



Although work discussed here has been restricted to a coupled geometry with a single extended interface, once such transfer process becomes reasonably controlled for multiparticle systems, one may envisage possible extensions to multiple connected closed-loop geometries (whether ring-shaped, race-track, or related), with the aim of deterministic transfer of winding numbers across a multiple-loop atomtronic architecture. Research into this promising direction is currently very active by the present authors.

Next, we discuss coupled persistent current dynamics in an alternative geometry of two transversally stacked ring-trap potentials connected by tunneling.

### C. Persistent currents in transversally coupled atomtronic circuits

Here, we briefly overview our recent findings<sup>402–404</sup> on dynamics of quantum vortices in two coupled vertically stacked toroidal condensates with persistent currents [see Fig. 31(a)].

In practice, the double-ring system with different angular momenta in its top and bottom parts may appear spontaneously as a result of cooling with different momenta,  $m_1$  and  $m_2$ , being frozen into the two rings after the transition into the BEC state, similar to spontaneous persistent current formation in coplanar coupled rings, described in Sec. VII B. We note that creation of ring currents in a double ring system by cooling<sup>421</sup> and instability of nonrotating tunnel coupled annular Bose–Einstein condensates<sup>422,423</sup> have been discussed in the literature.

The asymmetry of the density distribution in the top and bottom rings makes it possible to excite the vorticity also by applying a stirring laser beam, similar to the mechanism described in Sec. VII A. Generating a vortex in the lower-populated ring only, keeping the higher-populated one in the zero-vorticity state is illustrated in Fig. 31(b). A detailed analysis of the methods for persistent current generation in the system of coupled rings is under progress and will be published elsewhere.

In our recent work,<sup>402</sup> it was demonstrated that the azimuthal structure of the tunneling flows in double-ring system with topological charges  $m_1$ ,  $m_2$  implies formation of  $|m_1 - m_2|$  Josephson vortices, also known as rotational fluxons. The azimuthal structure of the tunneling flow [see the inset in Fig. 32(a)] implies zero net (integral) current through the junction for states, built of persistent currents with different topological charges in coupled rings ( $m_1 \neq m_2$ ). In particular, these include the case of opposite topological charges

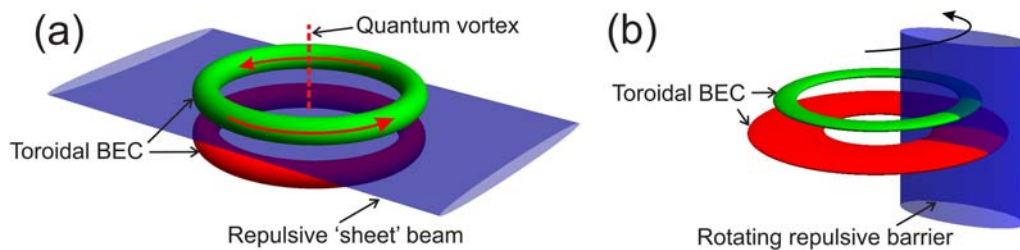
( $m_1 = -m_2$ )—considered in Ref. 424 and called “hybrid vortex soliton.” These structures host two different types of the vortices: vertical vortex lines and horizontal Josephson vortices. It turns out that the fluxons’ cores rotate and bend, following the action of the quench, i.e., the formation of tunnel junction with chemical potential difference. It was found in Refs. 402–404, as the barrier decreases, and the effective coupling between the rings, respectively, increases, the Josephson vortices accumulate more and more energy. When the persistent currents merge the relaxation process to new equilibrium state is driven by 3D dynamics of interacting Josephson vortices and vortex lines of the persistent currents [see, for example, Fig. 32(b)].

In our simulation of the merging rings, we have used the dissipative Gross–Pitaevskii equation in the following form:

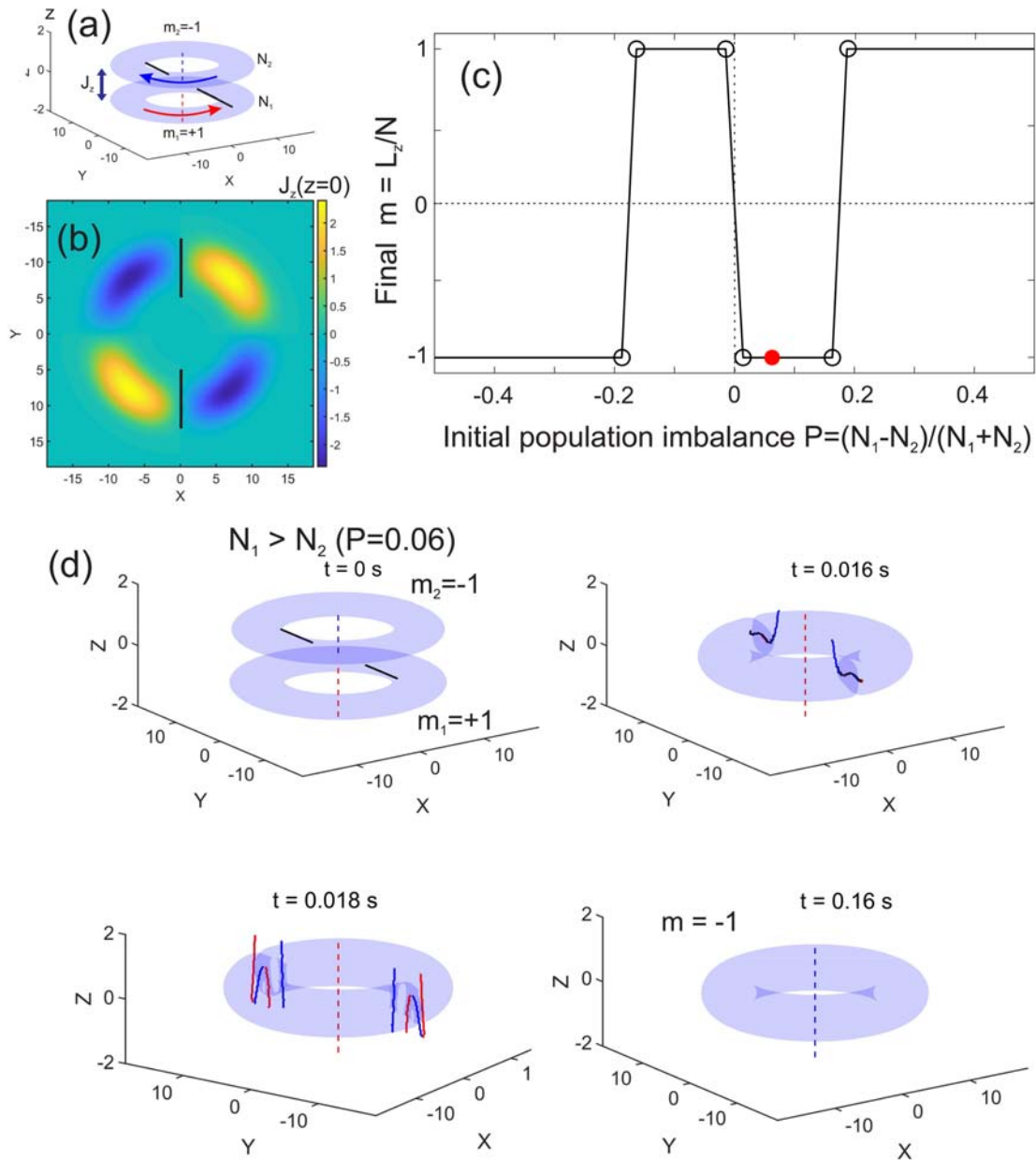
$$(i - \gamma)\hbar \frac{\partial \psi}{\partial t} = -\frac{\hbar^2}{2M} \nabla^2 \psi + V_{\text{ext}}(\mathbf{r}, t)\psi + g|\Psi|^2\psi - \mu\psi, \quad (60)$$

where  $g = 4\pi a_s \hbar^2 / M$  is the coupling strength,  $M$  is the atomic mass ( $M = 3.819 \times 10^{-26}$  kg for  $^{23}\text{Na}$  atoms),  $a_s$  is the  $s$ -wave scattering length (positive  $a_s = 2.75$  nm, corresponding to the self-repulsion in the same atomic species, is used below),  $\mu$  is the chemical potential of the equilibrium state, and  $\gamma \ll 1$  is a phenomenological dissipative parameter. This form of the dissipative GPE has been used extensively in previous studies of vortex dynamics (see, e.g., Refs. 382, 384, 385, 413, and 425). Note that main results of our work, concerning the role of the symmetry breaking in the interacting superfluids rings are not sensitive to the weak dissipative effects. We demonstrated in our works<sup>403,404</sup> that the symmetry of the system is the key feature explaining remarkable properties of the interacting quantized superflows. Certainly, the symmetry is in no way related to details of the dissipative terms. We have found that the subsequent relaxation process is determined by the initial stage of the evolution of the merging ring, in the course of several first microseconds after the barrier was switched off. Obviously, an effect of the weak dissipation on these fast processes is practically negligible. The dissipation plays a significant role in the course of subsequent temporal evolution of the condensate. In fact, in most experiments, *in situ* observation of the vortices is not possible, and only the final state can be analyzed after the completion of the relaxation. We include the dissipative effects in our model to investigate the final states of the merging superflows, which can be directly compared with expected experimental observations.

The relaxation of the merging rings is driven by substantially 3D nonlinear dynamics of the vortex lines corresponding to persistent



**FIG. 31.** (a) Schematics of the coupled ring-shaped condensates. Coaxial rings separated by repulsive potential, allowing investigation regime of tunneling coupling (long Josephson junction) and regime of merging rings (when the barrier is eliminated). (b) Schematics of preparation of the state with different angular momenta in double-ring system. Coaxial rings with different number of atoms are stirred by a rotating potential barrier. A persistent current is generated in a less populated ring (shown by green) while more populated toroidal condensate (shown by red) remains in nonrotating state.



**FIG. 32.** Coupled coaxial superfluid atomic circuits with counter propagating persistent currents. (a) Hybrid vortex stationary states with hidden vorticity. Vertical red and blue dashed lines designate cores of the counter-propagating persistent currents in the two rings. The cores of the Josephson vortices are indicated by solid black lines. (b)  $z$ -component of the corresponding tunnel-flow density distribution through the barrier,  $J_z(x, y, z = 0)$ . (c) The final value of the total angular momentum per particle  $L_z/N$  for the merging rings with initial vorticities  $(+1, -1)$  as a function of initial population imbalance  $P = (N_1 - N_2)/(N_1 + N_2)$ . (d) An example of evolution of the merging rings in oblate trapping potential. The barrier separating two rings is switched off at  $t > t_d = 0.015$  s, dissipative parameter  $\gamma = 0.03$ . Red (blue) lines indicate positions of the vortex (antivortex) core. The population of the bottom ring, with vortex  $m_1 = +1$ , is slightly larger than in the top one, with antivortex  $m_2 = -1$  [initial imbalance parameter,  $P = 0.06$ , is indicated by filled red circle in (c)]. The final state has  $m = -1$ . The symmetric drift of two diametrically opposite antivortices toward the central hole leads to subsequent annihilation of the central vortex and relaxation of the toroidal condensate into a final *antivortex*  $m = -1$  state, as described in Ref. 403.

currents and Josephson vortices, as illustrated in Fig. 32(b). It turns out that the final state of the condensate crucially depends on an initial population imbalance in the double-ring set, as well as on the shape of the 3D trapping potential, oblate or prolate.<sup>404</sup> In the

oblate (axially squeezed) configuration, a ring with nonzero angular momentum can impose its quantum state onto the originally non-rotating ring only above a well-defined critical value of the population imbalance.

It is apparent that two merging classical counter-propagating flows with zero total angular momentum evolve to the ground (nonrotating) state. Surprisingly, merging counter-rotating quantized flows in the axially-symmetric trap *never evolve toward the nonrotating ground state*, with  $L_z = 0$ , even for small imbalances,  $P \ll 1$  (see Fig. 32). It is particularly remarkable that the vorticity of the final state is imposed by the *less populated component* if  $P < P_{cr} \approx 0.1755$ , and by the stronger component only if  $P > P_{cr}$ . These counter intuitive properties of merging superflows are illustrated in Figs. 32(c) and 32(d). In this example, the number of atoms in the ring with topological charge  $m_1 = +1$  is moderately greater than number of atoms in the antivortex state ( $m_2 = -1$ ). The symmetric drift of two diametrically opposite antivortices toward the central hole leads to subsequent annihilation of the central vortex and relaxation of the toroidal condensate into a final antivortex state, i.e., the final topological charge of the merger is imposed by less populated ring as is seen in Fig. 32(d). A remarkable role of the symmetry of this system for vortex dynamics is investigated in Ref. 403.

Instead of the development of the classical Kelvin–Helmholtz instability at the interface of the merging persistent currents in a prolate potential trap, sufficiently elongated in the axial direction, we observe the formation of nonlinear robust hybrid vortex structures (as illustrated in Fig. 33 and explained in Ref. 404).

Thus, the ring-merging process and topological charge of the final state can be controlled by the perturbation of the trapping potential, specially adapted for the initiation of symmetry-breaking of the system, and by tuning of the initial population imbalance.<sup>403,404</sup>

#### D. Concluding remarks and outlook

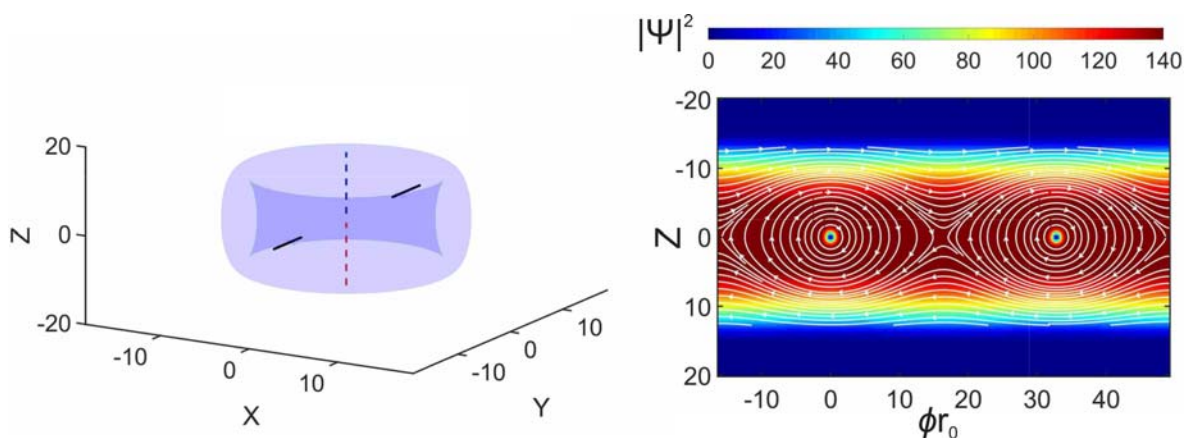
We have reviewed our current understanding of the spontaneous and controlled formation and stability of persistent currents in basic atomtronic circuits consisting of single or coupled ring-trap potentials and extended racetrack potentials. We have found that on-demand persistent flow can be created in a racetrack potential by stirring. The flow speed can be set to any value by adjusting the stir speed and/or the racetrack geometry. We discussed how persistent currents can also

be generated spontaneously after crossing the BEC phase transition. In coplanar geometries, we showed that the spontaneous generation of persistent flow is unaffected by the density overlap of the two rings, taking the first step in understanding ring–ring interactions and opening the possibility of many-ring arrays in the future. We have discussed our recent findings on dynamics of quantum vortices in two coupled vertically stacked toroidal condensates with persistent currents. It turns out that evolution of weakly coupled superfluid rings and merging quantized superflows with different topological charges is determined by complex dynamics of rotational Josephson vortices located between persistent currents.

The control over quantum topological excitations in such geometries offers an outstanding route to emerging quantum technologies with wide-ranging applications, such as topologically protected fault-tolerant quantum computation and quantum sensors for acceleration and rotation. These critically rely on minimizing decoherence and dissipation and optimizing the engineering of such components. For example, flexible sensor operation would require a rapid generation of the desired initial state, with further reduction in shot-to-shot atom number fluctuations crucial for sensor accuracy. Other areas where further theoretical and experimental work is needed (and currently well underway) include atomtronic transport, gatelike manipulation of quantum topological excitations, and readout mechanisms in atomtronic circuits.

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**FIG. 33.** A long-lived hybrid complex is produced by the evolution of merging strongly elongated toroidal condensates, as described in Ref. 404. Shown are 3D isosurface with constant condensate density (left) and maps of the distribution of the density (right) on the cylindrical surface at the radius corresponding peak density of the condensate,  $\phi$  being the angular coordinate.



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## VIII. PHASE SLIP DYNAMICS ACROSS JOSEPHSON JUNCTIONS

**A. Minguzzi, A. Pérez-Obiol, J. Polo, N. P. Proukakis, K. Xhani**

The phenomenon of superfluidity and its consequences can reveal itself in different ways in a quantum gas. One of the most remarkable manifestations of superfluidity is the frictionless motion of particles within the fluid, which is in direct relation with the macroscopic quantum phase coherence of the fluid. However, in certain circumstances, this frictionless motion can be broken with dissipation taking over. Phase slips represent one of the fundamental mechanisms leading to dissipation in superfluid systems.<sup>161,391,426–430</sup>

Phase slips correspond to jumps in the phase structure of the wavefunction of a quantum fluid. They can arise in dynamical superflow through a barrier in distinct manners, as summarized below: In the case of a coherent superfluid, dynamical flow through a barrier can trigger excitations that lead to phase slips. The form of such excitations depends on the dimensionality and the geometry of the system and can take the form of solitonic or vortex excitations, with associated acoustic emission. The presence of fluctuations in the system—whether of thermal, or quantum nature—creates an additional mechanism of “incoherent” phase slips, thus giving rise to richer dynamics. Atomtronic circuits typically consist of one (or more) Josephson junctions, embedded within a closed, typically ring-shaped and low-dimensional, geometry. Therefore, understanding phase slip processes can prove crucial for the development of quantum technologies and, in particular, Atomtronic devices.

Currents in ring geometries are ideal candidates for the study of superfluidity in interacting quantum gases. In particular, these currents are metastable states that can maintain the flow of particles even when no external field or force is applied. However, these metastable states can decay in different scenarios. The decay of current states in one dimensional rings corresponds to a sequence of phase slips associated with the loss of angular momentum by the system. Such a type of event may occur as triggered, e.g., by thermal fluctuations or other type of fluctuations, and in such case, they correspond to the aforementioned incoherent phase slips, as well as in quantum coherent manner, i.e., as an oscillation among different angular momentum states.<sup>431</sup> Examples of dissipative motion have been observed in hysteresis dynamics<sup>16,161,383,432,433</sup> where thermal activation plays an important role.<sup>434</sup>

In addition, phase slips can also be triggered by an external mechanism, for instance a weak link can catalyze the production of vortices at zero temperature<sup>435</sup> and solitons.<sup>163</sup> It has been shown that thermally activated phase slips can become dominant in the damping dynamics of some observables at relatively low temperatures.<sup>388,436</sup>

Recent studies have demonstrated the connection between the dissipative motion observed in Josephson systems and phase slips.<sup>162,430,437,438</sup> In this case, vortex nucleation can be triggered through the weak link producing the Josephsonlike junction<sup>29</sup> or through thermal activation, depending on the range of parameters.

In a similar case, the connection between low-energy excitations and dissipative motion was proven to be the main mechanism,<sup>439</sup> leading to the damping of Josephson oscillations.<sup>440</sup>

In this section, we present recent developments in the topic of phase slips and their role in the dissipative dynamics of observables such as population imbalance of Josephson systems and current dynamics in ring potentials. Special attention is given to the nonlinear excitations of the different systems, such as vortex rings and dark solitons. In Secs. VIII A–VIII E, we summarize different studies performed by the authors, which illustrate how phase slips can emerge in Atomtronic devices and isolated quantum systems.

Section VIII A is devoted to the study of nucleation of vortex rings in a weakly linked three-dimensional elongated superfluid. In Sec. VIII B, we consider the damping of Josephson oscillations in a one-dimensional (1D) strongly interacting Bose gas. Section VIII C is dedicated to the excitation spectrum of a 1D stirred Bose gas. Section VIII D focuses on the dynamical phase slips occurring in a phase imprinted Bose gas trapped in a ring potential. Finally, in Sec. VIII E, we present the conclusions and outlook, summarizing and discussing how phase slips play a crucial role on Atomtronic-based devices.

### A. Critical transport and vortex dynamics in a thin atomic Josephson junction

In this section, we give a detailed and intuitive picture of the emergence of phase-slips and dissipation across a single Josephson junction in a full three-dimensional (3D) atomtronic geometry. Although there exist phenomenological models which can account for dissipative effects in such a setting—such as the extension of the two-mode model of Josephson junctions<sup>441</sup> to include a damping term<sup>442–444</sup> or models based on the analogy to a resistively-shunted junction (RSJ) circuitry<sup>445</sup>—such models can only offer limited insight into the microscopic characterization of the observed dissipation. For a more complete discussion, this section, therefore, focuses on the case of a junction embedded within an elongated harmonically-trapped superfluid, as a paradigmatic example of the arising dynamics. Such a choice is based on the existence of a carefully characterized experiment,<sup>430,437</sup> detailed *ab initio* numerical analysis of which<sup>438,446</sup> has enabled not only qualitative connections to be made, but also facilitated direct links between microscopic and macroscopic observables and manifestations, directly relating these to the experimental observables. The discussion below is, thus, based on our recent works,<sup>438,446</sup> conducted at both zero and finite temperatures, which have fully analyzed all aspects of the arising microscopic dynamics.

The relevant experiment focused upon here was conducted in Florence,<sup>430,437</sup> in the context of an elongated <sup>6</sup>Li *fermionic* superfluid, separated by a thin Gaussian barrier induced by a focused laser beam located at  $x=0$  and of  $1/e^2$  width  $w \sim 4\xi \sim 2 \mu\text{m} \ll R_x \sim 110 \mu\text{m}$ , where  $\xi(R_x)$  denotes the superfluid healing length (axial system size). The experiment probed all regimes of values of  $(k_F a)^{-1}$ , where  $k_F$  denotes the Fermi wavevector and  $a$  the atomic *s*-wave scattering length. Although relevant and subtle differences were observed when transitioning from the BEC to the BCS superfluid regimes—mainly associated with different critical velocities and spatial extents due to the changing interaction dependence and the increasing importance of the fermionic degrees of freedom—the key underlying physical process leading to dissipation of superflow was found to be the same in all

regimes, as outlined below (see also Sec. XI for a discussion of transport and dissipation in ultracold Fermi gases).

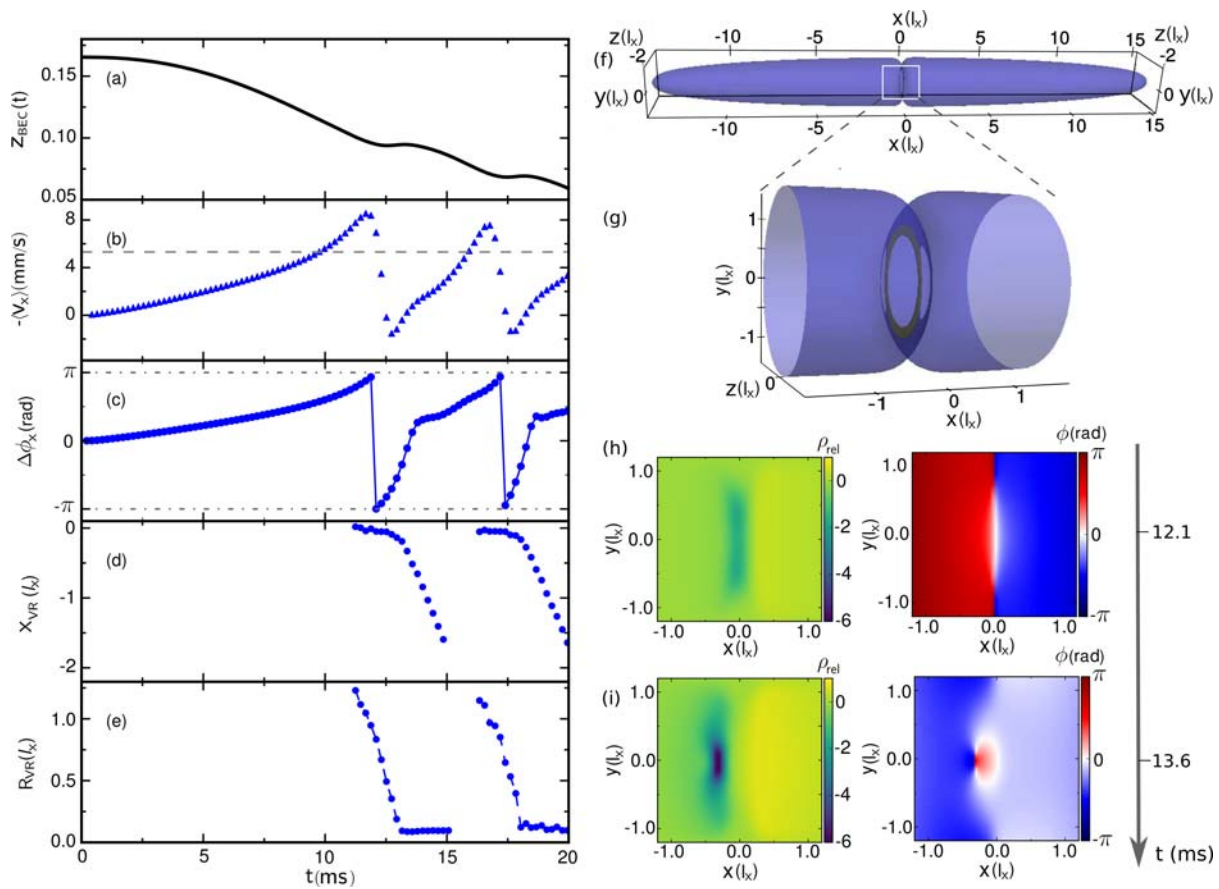
The presence of an initial population imbalance across the two wells separated by the barrier (initiated by moving the superfluid relative to the barrier at  $t = 0$ ) induced a neutral current flow in the negative  $x$ -direction, leading to the transfer of particles from the right to the left well. As expected, small values of initial population imbalance were found to lead to symmetric Josephson “plasma” oscillations about a zero population imbalance and associated oscillations in the relative phase.<sup>441,447</sup> Nonetheless, when the initial fractional population imbalance exceeded a critical value, the system population dynamics transitioned to a different regime. Based on earlier experiments with ultracold Josephson junctions,<sup>428,448,449</sup> one may have expected a transition to a so-called macroscopic quantum self-trapping regime, in which the population transferring oscillations proceed around a nonzero value of the population imbalance (i.e., one side of the junction always has a higher population than the other), and with a running phase,<sup>441,447</sup> the existence of such a regime has been argued to be related to the presence of a vortex ring in the barrier region, which annihilates within the weak-link region (but outside the region of observable condensate density).<sup>450,451</sup> Interestingly, a different regime was observed in the recent Florence experiment characterized by the observation of vortices in the superfluid bulk. Such a dissipative regime can arise when the emergent vortex ring acquires sufficient energy to overcome the barrier and penetrate the bulk superfluid. This is the regime analyzed in our recent work.<sup>438</sup> Specifically in our case—and, in particular, during the first part of the dynamical evolution when particle flow is still in a single direction—and for sufficiently large population imbalances, the very narrow nature of the barrier<sup>438</sup> was found to induce a local superfluid flow that accelerates in time and exceeds the local critical velocity for vortex excitation, which, in this geometry, takes the form of a vortex ring excitation. As a result, the presence of a Josephson current flow led to the generation of a vortex ring, associated with a phase jump of  $2\pi$  across the axial direction, and a flow which is no longer dissipationless. Such a phase slippage process is well documented in related contexts of Josephson junctions in superfluids and superconductors,<sup>426,427</sup> where it can be described in terms of *phenomenological* models.

Numerical analysis has shed more light on this process in the highly controlled environment of an ultracold atomic gas, modeled by the full 3D Gross–Pitaevskii equation describing the low-temperature regime of a weakly interacting condensate. Such analysis has been conducted based on the experimental parameters of Refs. 430 and 437 in the molecular BEC regime. Key findings reported in Ref. 438 are clearly summarized in Fig. 34. Interestingly, the observed dissipation arises as a combination of the transfer of incompressible kinetic energy from the particle flow to the vortex and the acoustic (phonon) emission (with the latter potentially amounting to a significant fraction of the total flow energy). More details about this phase-slip process and the associated acoustic emission can be found in Ref. 438. For completeness—before proceeding further with such characterization—we note that while such a dissipative regime was found in the limit of relatively low and narrow barriers [height  $\lesssim O(\mu)$ , and width four times the system healing length, for the parameter space probed in the Florence experiments<sup>430,437</sup>] a transition to a self-trapped regime also exists in the limit of broader and/or higher barriers; such analysis, providing a unified overview of the distinct dynamical regimes across a

single Josephson junction was presented in Ref. 446. The stability of this self-trapping regime is directly affected by thermal and quantum fluctuations<sup>452–457</sup> and by higher order tunneling processes,<sup>458</sup> which are known to gradually destroy such a state, eventually leading to oscillations about a zero population imbalance.

We proceed here by reviewing the phase-slip-related transition to the dissipative regime probed in the Florence experiments.<sup>430,437</sup> Consider the case when the fractional population imbalance  $z_{\text{BEC}}$  across the junction starts at a positive value (i.e., right well has higher population than left well): the induced Josephson dynamics leads to an (initial) superflow toward the left side of the junction [Fig. 34(a)], thus causing an initial decrease in the fractional population imbalance. However, as such an imbalance decreases at an increasing rate, implying an increase in the superfluid velocity (and corresponding superfluid current), there comes a point when the magnitude of the superfluid velocity exceeds some threshold value [loosely set by the mean speed of sound shown by the horizontal dashed line in Fig. 34(b)], acquires a temporally local maximum value, as a result of which it becomes energetically favorable for a vortex ring excitation to be generated at the barrier at  $x = 0$ . Such a process is associated with an abrupt jump of  $\sim 2\pi$  in the condensate relative phase, as shown in Fig. 34(c). The vortex ring generation instantaneously opposes the population transfer [leading to the flattening of  $z_{\text{BEC}}(t)$  visible in Fig. 34(a)] and can even lead to a reversal of the background superflow [i.e.,  $-\langle v_x \rangle$  changing sign in Fig. 34(b)] due to the additional “swirling” velocity of the induced vortex ring. The vortex ring, initially generated (as a “ghost” vortex) in the low density region outside the local transversal spatial extent of the BEC, remains initially within the axial barrier region  $x_{\text{VR}} \sim 0$  [Fig. 34(d)], shrinking transversally [Fig. 34(e)] and entering the Thomas–Fermi radius. After a short time, the accelerating vortex ring reaches the axial edge of the barrier (the superfluid density maximum is located at  $|x_{\text{VR}}| \sim 2w$ ) and starts traveling at a constant speed [linear part of decreasing  $x_{\text{VR}}(t)$ ], while maintaining its radius. A detailed instructive visualization of the overall superfluid geometry and the narrow nature of the barrier region can be found in Figs. 34(f)–34(i), which also displays the vortex ring generation and initial dynamics.

The long-term dynamics after the generation of a vortex ring from the decay of the superflow depends critically on the system parameters. If the initial population imbalance is relatively weak (but still above the required threshold for defect-inducing decay of superflow), a single vortex-ring may be generated, whose lifetime and subsequent dynamics outside the barrier depends on the value of the barrier height, as shown in.<sup>446</sup> However, in cases of larger initial population imbalance, after the first vortex ring has been generated and left the central region, the background superflow due to the remaining population imbalance, i.e., chemical potential difference, picks up its pace [around  $t \sim 13$  ms in Fig. 34(b)], until at some time later, when the previously generated vortex ring has already traveled a (potentially significant) axial distance from the barrier region, it once again exceeds the local critical speed and a second vortex ring is generated (around  $t \sim 16.3$  ms). This process can repeat itself, leading to even more vortex ring generation, until (due to the decreasing population imbalance) the background flow weakens to the point that it can no longer exceed the critical velocity. The resulting sawtoothlike profile of  $-\langle v_x \rangle$  [Fig. 34(b)] is typical of phase slippage phenomena seen in superfluid helium.<sup>427,459–461</sup> A generated vortex ring eventually decays either by



**FIG. 34.** Characterization of phase slip process and subsequent vortex ring dynamics in a weakly-interacting elongated 3D condensate: (a) Fractional population imbalance  $z_{\text{BEC}} = (N_R - N_L)/(N_R + N_L)$  across a thin Josephson junction against time. (b) Induced superfluid velocity along  $x$ , weighted over the transverse density in the  $x = 0$  plane. (c) Induced superfluid relative phase along  $x$ , evaluated at  $z = 0$ ,  $y = 0.81l_k$ , corresponding to the vortex ring semiaxis along the  $y$  direction at  $t = 12.1$  ms [see subplots (g)–(h)]. (d) and (e) Corresponding position  $x_{\text{VR}}$  and mean radius  $R_{\text{VR}}$  of the generated vortex rings as a function of time. (f) and (g) 3D density profile (density isosurface taken at 0.002 of maximum density) at  $t = 12.1$  ms, revealing the superfluid geometry and narrow barrier, along with a zoom-in to the highlighted central region [enclosed within the white rectangle in (f)] where the generated vortex ring (green near-circular structure) becomes clearly visible. (h) and (i) Corresponding planar ( $z = 0$ ) 2D snapshots of the condensate density (left) after subtracting the background density and scaled to its maximum value, and phase profiles (right) revealing the emergence and early dynamics of the first generated vortex ring at the indicated times. Spatial axes are plotted in terms of the harmonic oscillator length along the  $x$ -axis,  $l_k \approx 7.5 \mu\text{m}$ . Parameters for this figure:  $N_{\text{BEC}} = 60,000$  bosonic  $^6\text{Li}$  molecules,  $1/(k_F a) \approx 4.6$ ,  $z_{\text{BEC}}(t = 0) = 0.17$ ,  $\omega_x \approx 2\pi \times 15$  Hz,  $\omega_y \approx 2\pi \times 187$  Hz,  $\omega_z \approx 2\pi \times 148$  Hz (cigar-shaped trap), based on a double-well potential defined by  $V_{\text{trap}}(x, y, z) = (M/2)(\omega_x^2 x^2 + \omega_y^2 y^2 + \omega_z^2 z^2) + V_0 e^{-2x^2/w^2}$ , where  $M = 2m$  is the molecular mass,  $V_0 = 0.8\mu$  is the height of the Gaussian barrier and  $w \approx 2.0 \mu\text{m}$  is the barrier  $1/e^2$  width. Figure similar in spirit to individual plots shown (for other population imbalances) in Ref. 438.

shrinking into a rarefaction pulse during its axial propagation (as relevant for the case considered here), or by interacting with the transversal condensate boundaries as the transversal spatial extent decreases during its propagation toward the axial condensate edge.<sup>438</sup> In cases of high initial population imbalance, the time window between successive vortex ring generation events (depending on  $\hbar/\Delta\mu$  with  $\Delta\mu$  being the chemical potential difference between the two wells) can be shorter than the vortex ring lifetimes, thus allowing the coexistence of multiple sequentially generated vortex rings; such rings may further interact both indirectly (through their respective emitted acoustic waves), and directly (vortex–vortex interactions), potentially leading (for very high initial population imbalances) to reconnection processes, “leap-frogging” (sequential passage of one vortex ring through the other), or

even a “turbulentlike” regime (already discussed, for example, in 2D geometries<sup>462</sup>) this, in turn, leads to a highly complicated long-term dynamics of the population imbalance.

The experimental observations<sup>430,437</sup> are consistent with the picture described here. More concretely, the experiments led to the observation of one, or more, individual vortices, seen after removing the barrier (an added experimental complication required for imaging purposes), allowing the system to evolve and expand. This is consistent with the underlying picture described above, upon detailed consideration of the transversally asymmetric nature of the potential (which leads to excited, noncircular, vortex rings exhibiting Kelvin wave excitations), inherent fluctuations (which lead to asymmetric generation, propagation and decay of the vortex rings, and can thus explain the



presence of single/odd-number-of defects in experimental expansion images) and dynamical barrier removal (which is found to significantly extend the lifetime of generated vortex rings).<sup>438</sup>

Reference 438 also considered the role of temperature and thermal fluctuations by means of a self-consistent (“ZNG”) kinetic theory,<sup>413,416,417,463</sup> in which the condensate is described by a dissipative Gross-Pitaevskii equation which explicitly includes friction and collisional population transfer with the thermal cloud, the latter being treated by a quantum Boltzmann equation. This demonstrated that the presence of small thermal fluctuations does not significantly influence the above vortex generation process, although we have observed that a high enough temperature can in fact induce *additional* thermally-activated vortex rings; examples of the latter behavior in the one-dimensional context are discussed by means of a different finite-temperature model in Sec. VIII D below. Beyond the initial generation process, thermal effects were found<sup>438</sup> to have a significant role on the long-term vortex dynamics, where they act both to destabilize the otherwise symmetric motion of the vortex ring through the introduction of fluctuations, and to damp the motion through a mutual friction damping mechanism.<sup>464,472</sup>

The process that we have discussed here is generic and applies to any geometry and dimensionality, even though specific details will vary. For example, in the case of one-dimensional systems (see Secs. VIII B and VIII D), the underlying defects generated are dark solitons with the corresponding dynamics in ring traps of direct relevance to atomtronics discussed in Sec. VIII C.

Although the above discussion focused primarily on reviewing the emergence of dissipation across a single 3D junction embedded within a harmonic trap, it is pertinent to highlight here the important related work of quantum transport across an atomtronic “dumbbell” circuit, consisting of two reservoirs connected by a configurable linear channel of variable length and width:<sup>29</sup> this was both studied experimentally, and analyzed theoretically in the context of a superfluid acoustic model, a phase-slip model for the conductance, and via mean-field simulations. Such a work also highlighted the existence of Josephson plasma and dissipative dynamical regimes with the transition between such regimes observed (for given channel) at variable initial population imbalance. Moreover, Gauthier *et al.*<sup>29</sup> noted that the relative importance of sound and vortex energy as the origin of dissipative dynamics depends sensitively on the details of the geometry of the channel: for example, small channel widths which cannot support vortex dipoles, lead to the generation of unstable topological excitations which decay rapidly to compressible excitations. Thus, for small channels, they found the origin of dissipation to be sound-dominated. This is qualitatively consistent with the findings of Ref. 446, which characterized the dominant dissipation mechanism for sufficiently high barriers as the propagation of emitted sound waves.

## B. Bose–Josephson junction among two one-dimensional atomic gases: A quantum impurity problem

The one-dimensional (1D) geometry in ultracold Bose gases provides an ideal physical platform for the study of the quantum dynamical behavior of Bose–Josephson junctions, as the low dimensionality of these systems leads to the enhancement of quantum fluctuations and correlations. Recent experiments have realized and studied the 1D strongly interacting regime by using quasi-one-dimensional cigar-shaped

potentials in which the transverse motion of the particles is effectively frozen.<sup>359,440,466–470</sup> One-dimensional systems present features that clearly separate them from the higher dimensional ones, especially in the intermediate and strongly interacting limit where the motion of the particles is defined by its collective behavior. This collective motion is tightly connected to the low-energy excitation spectrum of the gas.

One dimensional systems are characterized by specific thermalization properties (e.g., to generalized Gibbs ensemble for integrable systems), which has been a topic of continuous interest.<sup>249,292,471,472</sup> Phase slips play a crucial role in the dissipative dynamics of quantities such as population imbalance in Josephson systems and current dynamics in ring potentials. For instance, phase slips are the only mean to change angular momentum in one-dimensional rings, as such rings cannot host vortices in the transverse direction, and they occur at the position of a localized barrier. In one-dimensional wires, phase slips occur when a soliton is formed or destroyed upon hitting the barrier giving rise to the junction.<sup>163</sup> Hence, Josephson junctions in one-dimensional systems are an appealing physical platform to investigate such damping phenomena and one of the simplest yet complete many-body systems displaying thermal and quantum phase slips.

Recent studies have investigated phase slips in different contexts: for instance in Ref. 473, they investigated two tunnel-coupled one-dimensional tubes placed side-by-side and characterized their low-energy physics described by unequal Luttinger liquids. Other approaches are also being investigated; e.g., in Ref. 474, they attribute the short-time evolution to multimode dephasing, while for longer times, they relate the relaxation to the nonlinear dynamics of the system.

Subsection VIII B 1 presents a study of the microscopic origin of phase slips in 1D bosonic Josephson junctions. Specifically, the analysis is performed in the strongly interacting regime by considering two weakly coupled one-dimensional wires in a head to tail configuration. The results and discussion presented here are adapted from.<sup>439</sup>

### 1. Model

The intermediate and large interaction regimes of a 1D Bose gas are difficult to treat, both numerically and analytically, due to the many-body character of the system. Using the Luttinger liquid (LL) theory<sup>475</sup> one can calculate the low-energy dynamical response of two strongly interacting one-dimensional bosonic fluids confined within an effective 1D waveguide of length  $L$ , tunnel-coupled through a weak link created by a barrier. In particular, in Ref. 439, we have studied the system’s response to a quench in the particle number difference between the two subsystems. By using a mode expansion of the density fluctuation and phase field operators from the LL theory and by defining the relative coordinates for the field operators, we identified the zero modes  $\hat{N}$  and  $\hat{\phi}_0$  as the relative population and phase differences between the two coupled wires, and  $\hat{Q}_\mu$  and  $\hat{P}_\mu$  as the relative coordinates for the excited modes. The resulting Hamiltonian reads

$$\hat{H}_T^{rel} = \frac{\hbar^2}{2ML^2} (\hat{N} - N_{ex})^2 - E_J \cos(\hat{\phi}_0) + \sum_{\mu \geq 1} \left[ \frac{1}{2M} \left( \hat{P}_\mu + \frac{\sqrt{2}\hbar}{L} (\hat{N} - N_{ex}) \right)^2 + \frac{1}{2} M \Omega_\mu^2 \hat{Q}_\mu^2 \right] \quad (61)$$

with effective mass  $M = \hbar K / 2\pi v L = K^2 m / 2\pi^2 N_0$ ,  $N_0$  being the average particle number in each tube and  $N_{ex} \ll N_0$  the excitation

imbalance, which may be tuned by a suitable choice of the initial conditions. It is worth mentioning that the center-of-mass coordinates are completely decoupled from the relative ones and they simply take the form of a harmonic oscillator, which can be readily diagonalized, not playing any role in the observable of interest.

We identify in Eq. (61) three terms: (i) a *quantum impurity particle* term corresponding to the two collective variables  $\hat{N}$  and  $\hat{\phi}_0$ , (ii) a bath of harmonic oscillators formed by the excited modes, and (iii) a coupling term  $\propto \hat{P}_\mu \hat{N}$ , obtained by expanding the second line of Eq. (61). Hamiltonian Eq. (61) has the same structure as that of the Caldeira–Leggett model.<sup>476–478</sup> However, it is important to remark that in our model, the bath of harmonic oscillators is intrinsic to the microscopic model, while in the Caldeira–Leggett model, it is phenomenologically introduced. The energy scales  $E_Q$  and  $E_J$  depend on interactions, the latter being renormalized by quantum fluctuations.<sup>406</sup> Concomitantly, the sound velocity and the Luttinger parameter vary with the interaction strength as described in Ref. 475. The first two terms of Eq. (61) correspond to the familiar Josephson Hamiltonian, where two regimes can be identified depending on the ratio of the Josephson energy,  $E_J$ , and kinetic energy,  $E_Q = \hbar^2/mL^2 = 2\Delta E/K$ , with  $\Delta E = \hbar\pi v/L$  being the level spacing among the phonon modes of the bath.

Case 1: Let us first consider the case  $E_J \gg E_Q$ , i.e., the Josephson potential term  $-E_J \cos(\hat{\phi}_0)$  dominates upon the kinetic energy term in Eq. (61). In particular, starting from an initial particle imbalance among the two wires, its dynamical evolution was obtained from the Heisenberg equations of motion,<sup>439</sup> leading to the quantum Langevin equation of motion with three dominant parameters: the Josephson frequency  $\omega_J = \sqrt{\omega_0^2 - \gamma^2}$  where  $\omega_0 = \sqrt{E_J E_Q}/\hbar$ , the memory-friction kernel  $\xi_N(t)$  whose large temperature properties are given by  $\langle \xi_N(t) \rangle = 0$  and  $\langle \xi_N(t) \xi_N(t') \rangle = 2E_J^2 k_B T / \hbar^2 M L v \delta(t - t')$  and a damping rate given by  $\gamma = \pi E_J / \hbar K$  (assuming a large frequency cutoff for the LL theory),

$$\ddot{\hat{N}} + \omega_0^2 \cos(\hat{\phi}_0) \hat{N} + \int_0^t dt' \gamma_N(t, t') \dot{\hat{N}}(t') = \xi_N(t). \quad (62)$$

Within this Josephson regime, two different behaviors depending on interactions exist. In the weakly interacting limit, where  $K \sim 1/\sqrt{g_{1D}}$  and  $v_s \sim \sqrt{g_{1D}}$  with  $g_{1D}$  being the 1D interaction strength, the predictions of the two-mode model in its small-oscillation limit are recovered, i.e.,  $E_Q \propto g_{1D}$  and  $\gamma/E_Q$  vanish for  $g_{1D} \rightarrow 0$ , yielding undamped Josephson oscillations. However, for strong interactions,  $E_Q$  increases, as it is related to the compressibility of the system, and  $E_J$  decreases, since it is renormalized by increasingly larger phase fluctuations. Thus, by inspecting the dimensionless damping rate  $\gamma_Q \equiv \gamma/\omega_0 = \pi\sqrt{E_J}/\sqrt{E_Q K}$ , one can predict that the Josephson oscillations will be more and more damped at increasing interactions.

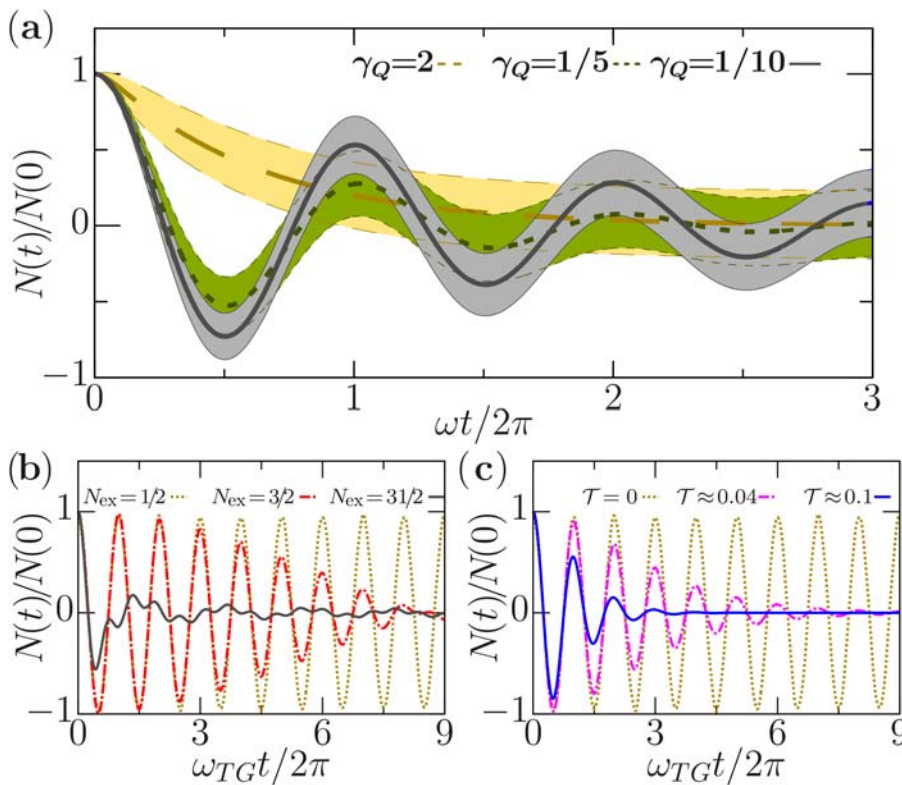
Case 2: In the  $E_J \ll E_Q$  limit, the phase is only weakly pinned and, therefore, it will display large fluctuations. In this regime, it is more convenient to use the Fock basis for the relative number. In this case, the energy levels of the quantum particle in Eq. (61) can be described as a function of the number of excitations  $N_{\text{ex}}$ , which now plays the role of quasi-momentum in the effective crystal, taking the form of parabolas  $\varepsilon_n(N_{\text{ex}}) = E_Q(n - N_{\text{ex}})^2/2$ , with  $\hat{N}|n\rangle = n|n\rangle$ . These parabolas present gaps of amplitude  $E_J$  opening at semi-integer values of  $N_{\text{ex}}$ . If we focus on the anticrossing points  $N_{\text{ex}} = \pm 1/2$ ,

$\pm 3/2, \dots$ , the system effectively behaves as a two-level model and the Josephson dynamics correspond to the Rabi oscillations of the quantum particle, with frequency  $E_J/\hbar$ . Note that the large value of  $E_Q$  fixes the scale of bath-modes level-spacing. This, creates a large gap between the level-spacing of the bath and that of the quantum particle,  $\hbar\omega$ . In the strongly interacting limit, there exists an exact solution for infinitely repulsive interactions, corresponding to  $K = 1$  in the LL theory, known as the Tonks–Girardeau (TG) regime.<sup>164,479,480</sup> In particular, in Fig. 35(b), we show that for a small initial imbalance, i.e.,  $N_{\text{ex}} = 1/2$ , undamped oscillations occur with a frequency  $\omega_{TG} = \varepsilon_{N+1} - \varepsilon_N$ . This is where the correspondence between the LL and TG regime can be made as  $E_J = \hbar\omega_{TG}$  and  $E_Q = \hbar^2\pi^2 N/mL^2$ . Hence, the oscillations observed in the exact solutions at small  $\delta V_0$  are the undamped Rabi oscillations of the quantum particle predicted by the LL model. For a larger imbalance, and thus beyond the low-energy description given by the LL theory, an effective damping appears due to the high energy excitations produced by the quench. The finite temperature regime was also addressed within the exact TG regime, as shown in Fig. 35(c) for small-imbalances. Unlike in the LL predictions, in the TG exact solution, damped oscillations were found. In order to pinpoint the origin of this damping, we computed the spectral function of the system at finite temperature.<sup>439</sup> We found that the exact spectral function contains multiple particle–hole excitations while the LL model assumes a linear excitation spectrum. In fact, we found that the exact spectral function also contains several low-energy excitations with frequencies of the order of  $E_J$ , which are associated with the presence of a finite barrier and are responsible for the observed damping.

In summary, our work showed that the LL model for two tunnel-coupled atomic gases can be mapped on a quantum impurity problem in the presence of a bath. The exact TG solution validates the frequency of the Josephson oscillations predicted in the LL model, and that the oscillations may in fact be damped by an intrinsic bath made out of low-energy excitations, but points out the existence of other modes that are beyond the LL model and that also provide damping of the excitations.

### C. Bose–Einstein condensate confined in a 1D ring stirred with a rotating delta link

Analyzing the spectrum of BECs trapped in ring settings is an important step toward understanding the generation and decay of supercurrents. The spectrum of a BEC in a 1D ring stirred by a rotating link can be first illustrated in the mean field limit, at zero temperature, and with a Dirac delta potential rotating at constant speed. Here, we follow these assumptions based on this section in Refs. 481 and 482. This approach has the advantage that the stationary solutions in the delta comoving frame are the ones of the free 1D GPE, and the effect of the moving potential is relegated to fixing specific boundary conditions. This is in contrast to models with finite width potentials,<sup>483–485</sup> and a generalization of a static point like impurity.<sup>10,486</sup> It allows for analytical expressions of the solitonic trains dragged by a rotating weak link and for the critical velocities at which the condensate becomes unstable and decays. The metastability of each excited state can be readily studied through a Bogoliubov analysis, and the hysteresis cycles observed in stirring experiments<sup>16</sup> can be qualitatively understood in terms of this model.



**FIG. 35.** (a) Relative number dynamics  $N(t)/N(0)$  (dimensionless) in two tunnel-coupled wires (LL approach) for various values of  $\gamma_Q = \gamma/\omega_Q$ . Stochastic noise uncertainties are indicated in shaded areas. (b) and (c) Relative-number oscillations (TG regime) following a quench of the step potential  $\delta V_0$  creating the initial imbalance: (b) at zero temperature for  $\delta V_0/E_F = 0.07$  (yellow-dotted line), 0.14 (magenta dashed line) and 0.72 (blue solid line), with  $E_F$  the Fermi energy; (c) at finite temperature for  $\delta V_0/E_F = 0.07$ . Reprinted with permission from Polo *et al.*, Phys. Rev. Lett. **121**, 090404 (2018). Copyright 2018, American Physical Society.<sup>439</sup>

## 1. Model

The stationary solutions are given by the condensate wave function,  $\phi(\theta)$ ,  $\theta \in [0, 2\pi)$ , and the corresponding chemical potential,  $\mu$ , in the delta comoving frame. Using natural units,  $\hbar = M = R = 1$ , with  $M$  the mass of the atoms and  $R$  the radius of the ring, the GPE and boundary conditions read

$$-\frac{1}{2}\phi''(\theta) + g|\phi(\theta)|^2\phi(\theta) = \mu\phi(\theta), \quad (63)$$

$$\phi(0) - e^{i2\pi\Omega}\phi(2\pi) = 0, \quad (64)$$

$$\phi'(0) - e^{i2\pi\Omega}\phi'(2\pi) = \alpha\phi(0), \quad (65)$$

where  $g > 0$  is the reduced 1D coupling, assumed to be repulsive,  $\alpha/2 > 0$  and  $\Omega$  the strength and velocity of the Dirac delta, and  $\phi$  is normalized to  $\int_0^{2\pi} d\theta |\phi(\theta)|^2 = 1$ . The spectrum is, thus, determined by three parameters,  $g$ ,  $\alpha$ , and  $\Omega$ . A general solution,  $\phi(\theta) = r(\theta)e^{i\beta(\theta)}$ , can be written in the closed form in terms of one of the 12 Jacobi functions.<sup>10</sup> These functions contain two free parameters, the elliptic modulus,  $m$ , which generalizes the trigonometric functions into the Jacobi ones, and a frequency  $k$ . Any set of values for  $k$  and  $m$  entails a solution that satisfies Eqs. (63)–(65) for a specific strength  $\alpha$ , velocity  $\Omega$ , and chemical potential  $\mu$ .

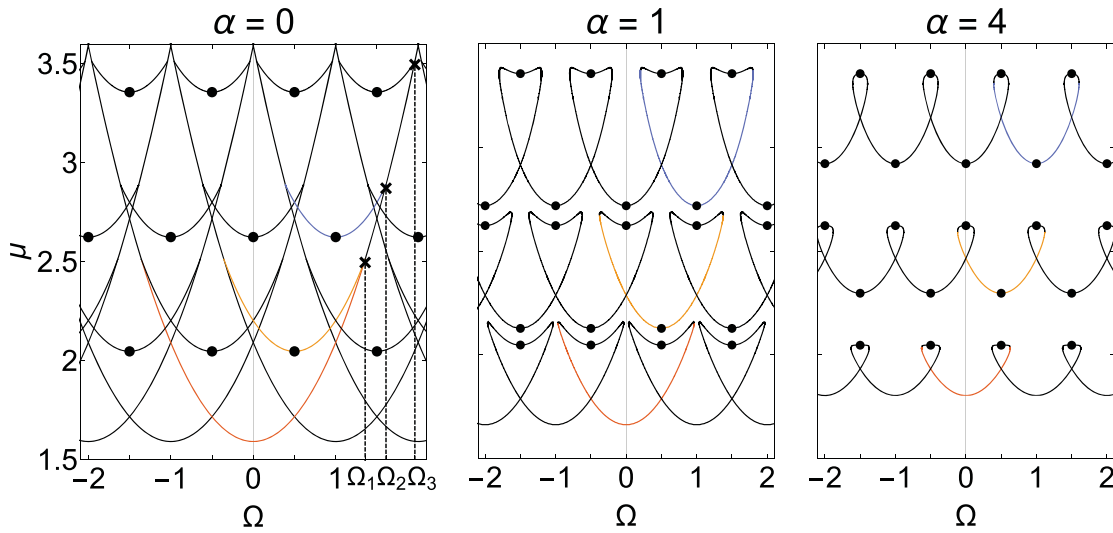
## 2. Spectrum

The free and stationary solutions,  $\Omega = \alpha = 0$ , consist in plane waves, real symmetry breaking solutions, and complex symmetry breaking solutions.<sup>487</sup> They correspond to vortex states, dark solitonic

trains with an even number of zeros, and gray solitonic trains. The latter is a generalization of the former two, plane waves representing the limit in which gray solitons become infinitely shallow, and dark solitons the limit in which the minima of gray solitons become zero. All these solutions are found by imposing a phase jump  $\beta(2\pi) - \beta(0) = 2\pi n$  with  $n$  an integer. If instead one constrains an arbitrary phase difference of  $2\pi\Omega$ , the obtained solitonic trains move at velocity  $\Omega$ —and are stationary in the frame of reference rotating at  $\Omega$ . The spectrum of stationary solutions from the point of view of an observer moving at  $\Omega$  is plotted in the left panel of Fig. 36. These solutions also include plane waves under a boost of  $\Omega$ . Dark solitonic trains with an even number of zeros comove with the condensate at  $\Omega = l$ , while trains with an odd number of zeros travel at  $\Omega = l + 1/2$ , where  $l$  is an integer. Waves moving at velocities departing from  $\Omega = l/2$  consist in gray solitonic trains, with shallower solitons the larger  $|\Omega - l/2|$ . At  $\Omega = l/2 \pm |\Omega_n - n/2|$  with  $\Omega_n = \sqrt{g/2\pi + n^2/4}$  and  $n$  indicating the number of dark solitons in the original train, the amplitudes become constant, and the gray soliton solutions merge into plane waves (parabolas in Fig. 36).

Once a barrier is created, the rotational symmetry is broken, and gray and dark solitonic train solutions are split into two. The energy spectrum, as observed in the Dirac delta comoving frame, is split into a set of swallowtail (ST) diagrams, see middle and right panels of Fig. 36. This looped structure implies that each wave train with a fixed number of dips in the density can be dragged only at a certain range of velocities. This range is limited by a pair of critical velocities, one  $\Omega < l/2$  and another  $\Omega > l/2$ , beyond which stationary solutions do not exist for the particular ST centered at  $\Omega = l/2$ . These pair of





**FIG. 36.** Lower part of the spectrum  $\mu(\Omega)$ , in the reference frame of a Dirac delta of strengths  $\alpha = 0, 1$ , and  $4$ , moving at constant velocities  $\Omega \in (-2, 2)$ , and where  $g = 10$ . Dots mark the velocities and energies of dark solitonic trains. Crosses in the left panel mark the velocities  $\Omega_n$  at which gray solitonic trains merge into the ground state for  $\Omega > 0$ . Reprinted with permission from Pérez Obiol and Cheon, Phys. Rev. E **101**, 022212 (2020). Copyright 2020, American Physical Society.<sup>481</sup>

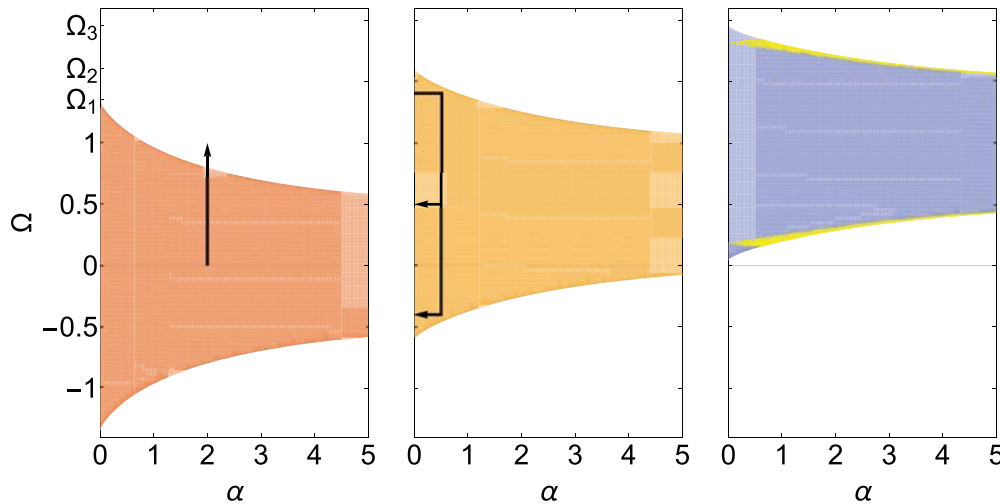
velocities are marked by the tips in each swallowtail, and depend on the magnitude of the weak link.

### 3. Metastability

The critical velocities define the regions in parameter space where stationary solutions exist, and the possible stirring protocols, or paths  $[\alpha(t), \Omega(t)]$  through which solitonic trains can be dragged, see Fig. 37 for a sample of these regions. Bogoliubov analysis show that the solutions corresponding to these regions are mostly stable under perturbations, while the solutions corresponding to top parts of swallowtails

are completely unstable. This metastability analysis does not qualitatively change for a wide range of nonlinearities ( $g = 1$  to  $g = 50$ ). In general, weaker atomic interactions and larger  $\alpha$  imply narrower ranges of velocities at which solitons can be dragged without dissipation. Moreover, for small  $g$ , the swallowtail loops become smaller, and at half-integer velocities, pairs of dark solitonic states are energetically very close to the ground states.

With the general features of the spectrum and its metastability laid out, one can devise adiabatic paths that avoid both, the critical lines delimiting the tips of the ST, and the metastable regions. Setting a weak link at zero velocity and then accelerating it, only produces



**FIG. 37.** Regions where stationary solutions exist corresponding to the bottom of the first three ST centered at  $\Omega = 0, \frac{1}{2}, 1$  and for  $g = 10$ . They are colored as the corresponding sections in Fig. 36. All these solutions are stable against perturbations except for the regions in yellow in the right plot. A path which explicitly leaves the stationary region is drawn on the left panel. The paths in the middle panel represent cycles to obtain a dark soliton and a vortex with one quantum of angular momentum. Reprinted with permission from Pérez Obiol and Cheon, Phys. Rev. E **101**, 022212 (2020). Copyright 2020, American Physical Society.<sup>481</sup>

currents of angular momentum  $J \lesssim 1$  before the condensate becomes unstable. If instead the weak link is set while rotating at a finite velocity  $\Omega_i \in (\Omega_n, \Omega_{n+1})$ ,  $n \geq 1$ , and then slowed down, any number of dark solitons or vortex states with any number of quanta of angular momentum can be obtained. As an example, cycles to obtain one dark soliton and a vortex are drawn in the middle panel of Fig. 37. The densities and phases corresponding to the vertices of this rectangular path are plotted in Fig. 38.

One can also devise paths that explicitly cross the critical line separating the stable and unstable regions. One such path is schematically plotted in the left panel of Fig. 37, where a weak link is set in the condensate at rest and then accelerated passed the critical velocity. In this case, one can expect the condensate to enter an unstable state and decay to the immediate lower state, corresponding to the lower branch of the swallowtail diagram. If then the opposite path is taken, in which the weak link is slowed down, the previous states are not recovered, and the condensate is left with an increased angular momentum, producing hysteresis. To recover the initial state, the weak link velocity has to be further decreased to reach the other critical point, so that the condensate decays again and the hysteresis cycle can be closed. Therefore, the swallowtails and critical velocities also provide a basis model to understand the hysteresis cycles observed in experiments. In particular, in Ref. 16, the cycle widths and critical velocities decrease as stronger weak links are rotated. This qualitatively agrees with the spectrum studied here, where the hysteresis widths  $\Delta\Omega$  are defined by the widths of the swallowtails, and decrease as weaker interactions  $g$  or larger link strengths  $\alpha$  are considered. Moreover, the model also

predicts hysteresis cycles coupling states with different angular momentum, corresponding to paths along upper swallowtail diagrams. On the other hand, these downward swallowtails are the characteristic of repulsive interactions, and it can be shown that such hysteresis cycles cannot happen for attractive interactions.<sup>482</sup>

The above stirring mechanisms, deduced analytically from the spectrum, are corroborated by numerical simulations of the time-dependent GPE where a Gaussian potential is rotated, instead of a Dirac delta. These simulations also allow one to test the stability of stirring protocols involving more excited states not studied in the analytical case. Indeed, for Gaussian widths of about 2% of the ring perimeter, dark solitonic trains with various zeros and vortices with up to a few quanta of angular momenta are produced following the protocols provided by the model with a rotating Dirac delta. Similarly, the condensate is able to sustain stable solutions when stirred by a Gaussian link up to a certain velocity. This critical velocity decreases with the Gaussian height, as expected from the regions of stationary solutions in Fig. 37.

This model offers a new approach to study metastability and vortex and soliton nucleation in ring condensates with a rotating weak link. The processes described in this section can also be understood in terms of a rotating trap and a fixed weak link or defect. Analytical expressions for the dragged solitonic trains and critical velocities allow to study ground states as well as excited states, and to understand how they are coupled among them. At half-integer velocities and for strong enough weak links dark solitonic states with zero current could be easily excited from the ground states. They could also be produced if a

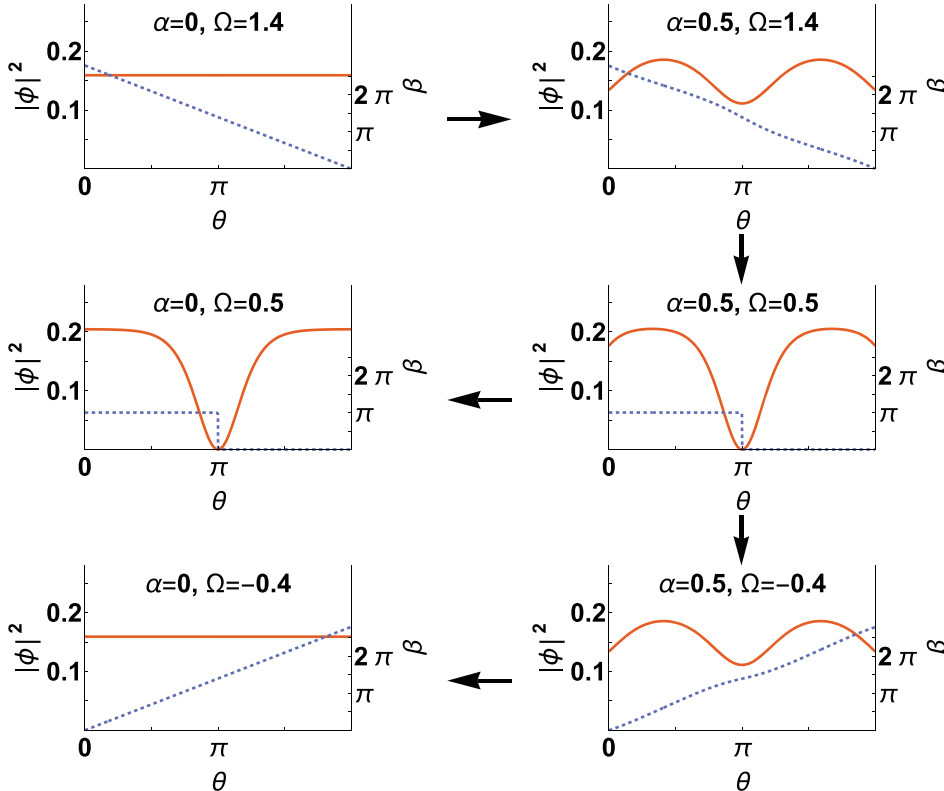


FIG. 38. Densities (solid lines) and phases (dashed lines) in the Dirac delta comoving frame corresponding to the key points of the dark solitons and vortex cycles in Fig. 37 (where  $g = 10$ ). The first steps in both of them consist in setting a delta on the ground state (top left plot) while rotating at  $\Omega = 1.4$ , and then lowering its velocity to  $\Omega = \frac{1}{2}$  (top and middle right plots). At this point, one can unset the delta and obtain a dark soliton, or further decrease the velocity down to  $\Omega = -0.4$ , and then unset the delta, in which case the state of one vortex is reached. Reprinted with permission from Pérez Obiol and Cheon, Phys. Rev. E **101**, 022212 (2020). Copyright 2020, American Physical Society.<sup>481</sup>

defect appears in a rotating condensate, which is then slowed down. Bogoliubov analysis indicate that the spectrum is roughly divided in stable and unstable regions (bottom and top parts of swallowtails). At the same time, the main features of hysteresis cycles can be qualitatively understood in terms of the swallowtail structure of the spectrum. This analysis complements phase slip mechanisms studied in 2D and 3D traps, where richer dynamics involving vortex excitation is possible, see Sec. VIII A, and numerical works in 1D where either stronger interaction regimes are studied, see Sec. VIII B, or explicit thermal activation is considered, as discussed next in Sec. VIII D.

#### D. Thermal and quantum phase slips in a one-dimensional Bose gas on a ring

This section's results and discussions are adapted from Ref. 162 and give an example of a one-dimensional quantum system where both coherent and incoherent phase slips appear at different regimes of bosonic interactions and temperatures. The section focuses on phase slips between different angular momentum states occurring in a one-dimensional Bose gas trapped in a ring potential. The current is induced through phase imprinting<sup>150,488,489</sup> and due to the dimensionality of the system vortex nucleation is forbidden within the ring. Therefore, one-dimensional phase slips require the existence of a different microscopic mechanism.

Previous studies have investigated the origin of phase slips and how these can lead to a decay of the superfluid current in different scenarios. For instance, in Ref. 436, they introduced a shallow lattice to trigger such phase slips and investigated the system using a combination of techniques based on the mean field Gross-Pitaevskii equation, including a Bogoliubov analysis as well as phenomenological noise and dissipation term in the mean field description. In a more recent work, it has been shown that phase slips can also be driven by acoustic waves in higher dimensional system,<sup>490</sup> which indicates that low energy excitations can be one of the triggering mechanisms of phase slips.

The work presented below covers all interaction range using different approaches and models. It also considers an experimentally realistic scenario where the currents are introduced through phase imprinting and takes into account how phase slips are originated in each regime. Therefore, it provides a good example of the current techniques and observations found currently in the field.

#### 1. Model and methods

As presentend in Ref. 162, we consider  $N$  bosons of mass  $m$  with repulsive contact interactions on a ring of circumference  $L$  with periodic boundary conditions. The coupling between different angular momentum states is triggered by the presence of a barrier, a procedure analogous to the experimental implementations of phase imprinting.<sup>150,488,489</sup>

In order to investigate phase-slips in the system, let us start from the equilibrium state  $\Psi_0$  in which a static barrier is present, and then induce a quench in the many-body wavefunction such that  $\Psi_0(x_1, \dots, x_N) \rightarrow \Psi_1(x_1, \dots, x_N) = \Psi_0 \times e^{i2\pi\ell \sum_j x_j/L}$ . The current dynamics is then obtained from

$$J(t) = -i \frac{\hbar}{2mN} \int_0^L \frac{dx}{L} \langle \Psi^\dagger \partial_x \Psi - (\partial_x \Psi^\dagger) \Psi \rangle. \quad (66)$$

The dynamical response is found using different methods depending on the interaction strength,  $\gamma = mg/\hbar^2 n$ , and temperature regimes: (i) the Gross-Pitaevskii equation (GPE) and analytical two-mode model adapted from Ref. 441 at  $T=0$  for a weakly interacting gas ( $\gamma \ll 1$ ); (ii) the projected Gross-Pitaevskii equation (PGPE) at  $T>0$  and  $\gamma \ll 1$ <sup>414,417,491</sup> and (iii) the time-dependent Bose-Fermi mapping at  $\gamma \gg 1$ , describing the infinitely strong interaction Tonks-Girardeau (TG) limit for the entire temperature range.<sup>164,479,480</sup> Throughout this section, a quench inducing a circulation  $\ell = 1$  is considered, although the results can be generalized to larger circulations. Depending on the model, two types of barriers are considered: a delta potential  $V(x) = \alpha\delta(x)$ , for which analytical results can be obtained, and a Gaussian potential  $V(x) = V_0 \exp(-x^2/2w^2)$ , which is more realistic from the experimental point of view. Both cases are compared using a dimensionless parameter for the barrier strength:  $\lambda_{GP} = V_0/\mu_0$  for weak interactions with  $\mu_0 = gn$  being the chemical potential of the homogeneous annular gas, and  $\lambda_{TG} = V_b/E_F$  for strong interactions, with  $V_b = \alpha n$  being the energy associated with the barrier and  $E_F = \hbar^2 n^2 \pi^2 / 2m$  the Fermi energy.

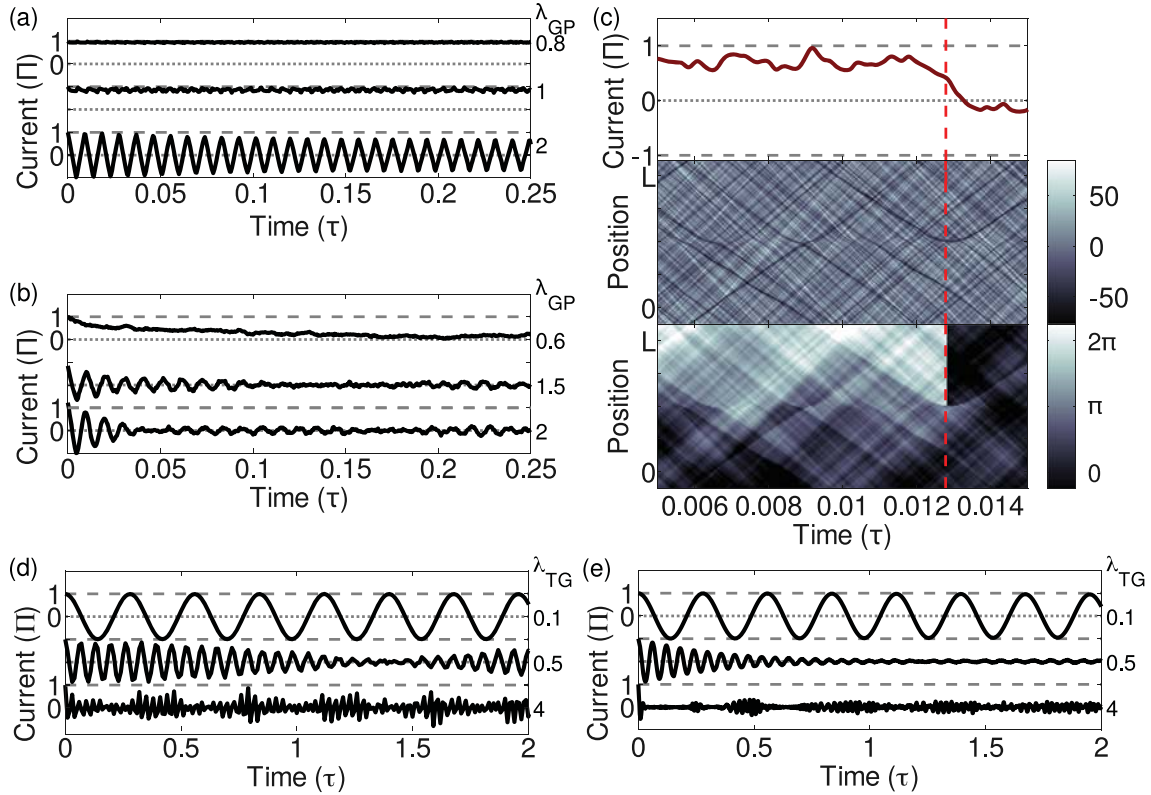
#### 2. Discussion

Within the considered system, the current dynamics depends on interaction and temperature regimes.<sup>162</sup> Figures 39(a)–39(c) show the results in the weakly interacting regime. At zero temperature, the current remains very close to the initial quenched circulating state for weak to moderate barriers, up to  $\lambda_{GP} \sim 1$ . Above this critical value, a fast decay of the current appears, followed by oscillations around the 0 value. This behavior is found to correspond to the transition of the currents from self-trapping to Josephson oscillations, in analogy to the well known Josephson effect for particle imbalance predicted in Ref. 441.

For temperature  $T = \mu_0/k_B$ ,<sup>492</sup> the dynamics of the current are quite different. At low barriers, i.e.,  $\lambda_{GP} \leq 0.5$ , we observe an exponential decay of the current, while for larger barriers, one observes damped oscillations. In this regime, thermal phase slips occur deterministically at the position of the barrier where the density vanishes. In order to elucidate the mechanisms behind the current decay, Fig. 39(c) shows a single classical field trajectory, showing many spontaneous thermal gray solitons.<sup>493</sup> While most of the solitons present a small density dip, thus being fast and transmitted through the barrier,<sup>494</sup> notice that the current undergoes discrete jumps each time a soliton reflects on the barrier. In this case, the density profile vanishes when the soliton reaches zero velocity, allowing for a phase slip to occur. This corresponds to the adiabatic process indicated by the dashed red line in Fig. 39(c). The observed exponential decay of the average current can be understood as an intrinsically stochastic process occurring when the barrier couples the soliton dynamics to the long wavelength sound excitations.<sup>494</sup>

The strongly interacting regime  $\gamma \gg 1$ , where the classical picture does not apply, is described using the exact Tonks-Girardeau solution. We show that the dynamics of the current microscopically corresponds to quantum coherent oscillations between different angular momentum states. At zero temperature, in contrast to the weakly





**FIG. 39.** (a)–(c) Classical field simulations of the quench dynamics in the mean-field regime for  $\gamma = 0.02$ . (a) Average current per particle [solid lines, unit:  $I = \hbar/(Nm)$ ] as a function of time (unit:  $\tau = mL^2/\hbar$ ) at  $T=0$ . Top to bottom:  $\lambda_{GP} = \{0.8, 1, 1.05, 2\}$ . (b) Current at  $T = \mu_0/k_B$  for barrier strengths  $\lambda_{GP} = \{0.6, 0.9, 1.5, 2\}$ . (c) Zoom on a single classical field trajectory at  $T = \mu_0/k_B$  and  $\lambda_{GP} = 0.6$ , illustrating a phase slip. This consists in a jump in the current (top panel), corresponding to the reflection of a slow soliton on the barrier, also visible in the density deviation map (middle panel) and appearing as a singularity in the phase profile (bottom panel). (d) and (e) Exact solutions in the Tonks-Girardeau regime. (d) Average current per particle versus time after the quench for  $N=23$ , at  $T=0$ , for barrier strength  $\lambda_{TG} = \{0.1, 0.5, 1, 4\}$ . (e) Current at  $T = E_F/k_B$  (solid lines) for  $\lambda_{TG} = \{0.1, 0.5, 1, 4\}$  from top to bottom. (a)–(e) Horizontal dotted (dashed) lines indicate the values for  $J=0$  ( $\pm 1$ ). Reprinted with permission from Polo *et al.*, Phys. Rev. Lett. **123**, 195301 (2019). Copyright 2019, American Physical Society.<sup>162</sup>

interacting regime, it can be seen that for weak barriers ( $\lambda_{TG} \ll 1$ ), there is no self-trapping [see Fig. 39(d)]. Rather, the current undergoes Rabi-like oscillations. These oscillations correspond to coherent quantum phase slips due to backscattering induced by the barrier, which breaks the rotation symmetry thus coupling different angular momentum states.<sup>406,431</sup> Microscopically, it corresponds to dynamical processes involving the whole Fermi sphere, i.e., multiple-particle hole excitations where each particle coherently undergoes oscillations of angular momentum from  $L_z = \hbar$  to  $L_z = -\hbar$ . At increasing barrier strength, an envelope appears on top of the current oscillations, degrading the Rabi oscillations. This envelope originates from the population of higher-energy modes, each transition being characterized by a different frequency (see Ref. 162) leading to a mode-mode coupling and dephasing, and more complex current oscillations. At finite temperatures, the quench dynamics of the current involves high-energy excitations with an amplitude weighted by the Fermi distribution.<sup>162</sup> The resulting dynamics corresponds to an effective damping of the current oscillations with an exponential decay [see Fig. 39(e)] due to the effect of incoherent phase slips. The revivals observed for large barriers at zero temperature are highly suppressed due to the thermal

excitations. We identify the oscillation frequency as Josephson oscillations in which at increasing barrier strength, the frequency crosses over from a Rabi-like regime with  $\omega = \pi^2 N \lambda_{TG}$  to a Josephson-like regime with  $\omega \propto \sqrt{\lambda_{TG}}$ . This is in agreement with the predictions of the low-energy Luttinger liquid theory (see Ref. 439 and Sec. VIII B above).

In summary, in this section, we have presented a study of the dynamics of a one-dimensional ring pierced by a localized barrier, following a phase imprinting. From a static point of view, a localized barrier can lead to solitonic excitations as seen in Sec. VIII C. However, these can also be thermally activated or created dynamically by quenching a current in the system. Within the mean field regime, the self-trapping behavior of the current prevents coherent phase slips, but at finite temperatures, incoherent phase slips are observed. Their microscopic origin is related to the coupling between the soliton dynamics and the long wavelength sound excitations, which are intrinsically stochastic, leading to an exponential decay of the average current. A similar microscopic behavior is found in higher dimensions; however, the excitations take other specific forms such as vortex rings as seen in Sec. VIII A. When considering the strongly interacting

regime, coherent phase slips dominate the dynamics. Finally, at finite temperatures, incoherent dynamics appears due to thermally occupied high-energy excitations that lead to an average decay of the current.

### E. Concluding remarks and outlook

We have discussed different regimes of ultracold atomic gases in which phase slips play a crucial role on the dissipative motion of certain macroscopic observables and discussed their connection to low-lying and macroscopic excitations in such systems. Understanding these processes in detail is crucial to harnessing future atomtronic applications.

To understand the microscopic origin of such mechanisms, we have considered the diverse settings of 3D harmonic traps and 1D systems in harmonic and ring traps with weak and strong interactions. We have also considered the effect of thermal fluctuations on such dynamics.

In the context of a weakly interacting harmonically confined 3D ultracold quantum gas with a weak link creating a Josephson-like junction, the phase slips are related to the generation of vortex rings and associated sound emission with increasing population imbalances leading to sequential ring generation, even potentially opening up an avenue for a turbulentlike regime. Our analysis was performed for a bosonic system, but the relevant experiment is actually performed across different superfluid regimes of an ultracold fermionic gas. As such, our results only strictly apply to the BEC side, and numerous interesting open questions remain on how the explicit nature of the fermionic statistics affects this picture as one moves toward the unitary and BCS superfluid regimes.<sup>430,437,495</sup>

In a 1D strongly correlated Bose gas, it is low-energy excitations within the bulk that provide the underlying mechanism leading to the dissipative motion across the junction. Although in this regime, we also observe that damping of large particle imbalances proceeds through higher-energy modes. On the other hand, in 1D ring potentials, the relevant excitations leading to the decay of the current at weak interactions are dark solitons, although low-energy excitations are also involved in the decay mechanism. In our studied regime, dark solitons were thermally activated; however, they can also have other origins, e.g., being triggered by the presence of an impurity. In that case, the specific soliton or solitonic train generated depends on the size of the impurity and its velocity relative to the current. The microscopic origin of the phase slippage is then related to the coupling between the soliton dynamics and the long wavelength sound excitations (which are intrinsically stochastic). We note that, while the 1D systems considered in Secs. VIII B–VIII D allow us to identify the microscopic origin of excitations, the barriers and topologies considered in these microscopic theories are significantly simplified ones, and extending this work to more realistic scenarios could bring new insight regarding the energy scales at which the damping of oscillations occurs. Indeed, present-day experiments use finite width barriers and external confinements to trap strongly correlated atoms, which could influence the damping. Moreover, the results shown in Sec. VIII B rely on the low-energy theory given by the Luttinger liquid model. Beyond the low-energy model, one should approach the problem numerically. However, numerical simulations of strongly correlated systems are highly complex. Therefore, developments in this field could prove of great importance in corroborating and extending the dynamics of strongly correlated Josephson coupled systems for strong

quenches. Finally, we note that the spectrum and role of impurities presented in Sec. VIII C and the microscopic mechanism leading to damping can notably depend on the dimensionality.

One of the main challenges for quantum technologies is to control the system's quantum state while maintaining its quantum coherence for longer times.<sup>2</sup> Thus, reducing dissipative motion becomes crucial for the development of atomtronic devices. Moreover, controlling and understanding the mechanisms involving coupling to low-energy excitations can also lead to a reduction of this dissipation. In addition, the initial quantum state can also be of consequence to the system's final stability, as the projection to high energy excitations can lead to complex damped dynamics. From these results, we can draw some insight regarding future directions for improving and reducing dissipative behavior. Integrable or quasi-integrable systems, in which many conserved quantities exist compared to the system's degrees of freedom, have been shown to present a long-lived coherence and dissipation-free dynamics.<sup>249,292,300</sup> Also, several recent studies have focused on topologically protected states<sup>196,497</sup> as the main building blocks for future atomtronic devices, as these states prove very robust against perturbations.

### ACKNOWLEDGMENTS

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### IX. ATOMTRONICS ENABLED QUANTUM DEVICES AND SENSORS

**D. Anderson, V. Ahufinger, A. S. Arnold, G. Birkel, M. Boshier, S. A. Gardiner, B. M. Garraway, J. Mompart**

In this section, we will discuss some example cases where the atomtronics approach is leading to novel components (which may be part of a larger device) and applications such as rotation sensing and magnetic sensing.

### A. Diodes, transistors, and other discrete components

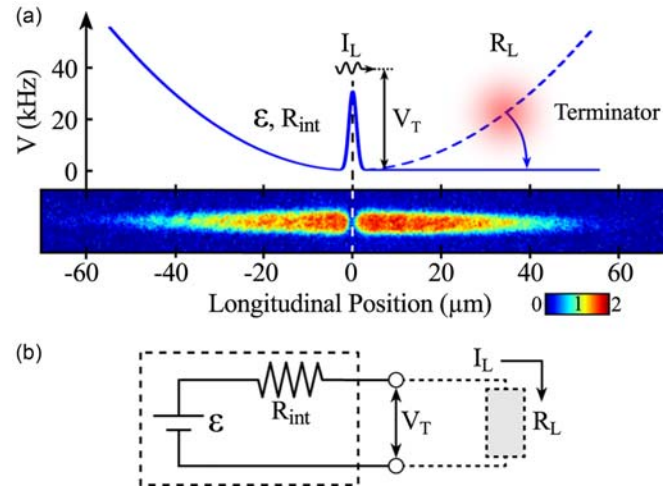
The terminology *Atomtronics* suggests, correctly, but not exclusively, an analogy between circuits for atomic matter and those based on standard *electronics*. The flow of electrons in an electric circuit can be considered in a way analogous to the flow of cold neutral atoms in an atomtronic circuit. (At present, there is some flexibility in this interpretation.) Important questions concern how to confine the atoms to a circuit, how to control them, and what devices and applications can arise.

It would be a gross over-simplification to suggest that an atomtronic circuit should merely mimic an electronic circuit. While this *might* be the case, it is by no means essential, and indeed, it is very much intended that future atomtronic systems go beyond analogs of standard electronic circuits. To be specific: Although we may start with these basic analogs, the future hope is for devices that use the properties of matter-wave coherence and other quantum properties of matter to go beyond these direct analogs, and to even create devices with no electronic counterpart because of the unique properties of the quantum physics of matter. Meanwhile, however, in this first part of Sec. IX, we will explore the progress made in formulating and implementing discrete atomtronic components that are similar to electronic ones.

The basic electronic elements are often regarded to be batteries, resistors, capacitors, diodes, transistors, and the like. If we start with the battery, in the atomtronic world, it can be regarded as a reservoir of atoms. Clearly, that is too simplistic and not enough on its own. So first steps are to involve atoms contained in a reservoir and allowed to flow out of that reservoir into a more complex circuit or at least into another reservoir as in cases of two-terminal flow.<sup>498</sup> The current of neutral atoms is driven by the difference in the chemical potential between two reservoirs, typically implemented by the two sides of a barrier, in a way that is analogous to Ohm's law in conventional circuit theory. The battery is later intended to supply power to an atomtronic circuit via the transport of cold atoms.

Such a battery was demonstrated in Ref. 499, which was complemented by the respective theoretical description in Ref. 500. In this “battery” experiment, a relatively large confining potential for ultracold atoms is divided into two parts by introducing a spatially narrow beam of blue-detuned light acting as a barrier [see Fig. 40(a)]. Atoms can be confined on one side of the combined potential, the other side, or both. During the experiment discussed here, atoms are initially loaded on one side [the left side in Fig. 40(a)] and then allowed to flow to the right side as controlled by the sharp barrier which essentially controls the “resistance.” This flow can be classical, or quantum, depending on the temperature of the loaded atoms. [Tunneling does not play a role in Fig. 40(a) if the main contribution to flow is from over-the-barrier atoms and if the left well rethermalizes sufficiently to maintain the distribution.] To prevent the reflection of atoms from the second well, a “terminator” is added to the system by means of an optical beam that pumps atoms in the second well into untrapped states, such that they are lost from the system. This terminator also represents a load-matched impedance [see Fig. 40(b)] in the analogous electric circuit.

A cold atom system with ballistic atoms has been used to demonstrate the dynamics of a linear *RLC* circuit.<sup>501</sup> In this case, a light sheet trap is modified with additional dipole beams to create 2D confinement of the atoms: two reservoirs are generated that are joined by a



**Fig. 40.** An atomtronic battery. (a) The confining fields for the atoms are formed using magnetic and optical potentials. The image of atoms demonstrates the situation when they fully occupy the system with the terminator off. (b) An equivalent circuit for the atomtronic battery. Reprinted with permission from Caliga *et al.*, New J. Phys. **19**, 013036 (2017). Copyright 2017, Author(s) licensed under a Creative Commons Attribution 3.0 License.

narrow link. These “capacitor”-like systems are charged by loading atoms into one of the reservoirs. Subsequently the flow of atoms through the narrow channel discharges the capacitor. The channel possesses a finite resistance and appears to have inductance as well.<sup>501</sup>

In analogy to solid state materials modifying the electronic wave function, optical lattices offer band-gap structures for cold atoms (in this context, see, e.g., Ref. 502). These allow the creation of diodes and transistors by changing the base-line potential of the lattice across a discontinuity or junction in just the same way as for semiconductors across a NP- or PN-type junction.<sup>11,503</sup> However, it is interesting to note that the atomtronic diode can display its functionality with just a few lattice sites (i.e., with just a few potential minima) and the atomtronic transistor is proposed to be functional with just three potential wells.<sup>30</sup> It can be constructed in the same way as the battery experiment discussed above,<sup>499</sup> but with an additional blue-detuned dipole beam adding one additional barrier to the passage of the atoms. The transmission of atoms through the double-barrier system is now dependent on the chemical potential between the two barriers,<sup>172,504</sup> giving a transistorlike behavior with “source,” “gate,” and “drain” assigned to the three regions around the barriers. Furthermore, by cascading transistor junctions, i.e., by adjusting the sets of lattice potentials, a logic gate (AND gate) has been proposed consisting of just five lattice sites.<sup>11</sup>

One direction for future extensions is the development of more exotic circuit elements. For example, asymmetric double well potentials made by optical dipole beams have been used to create a Josephson junction for an atomtronic system.<sup>505,506</sup> In the direction of increasing complexity of circuits, for example, the proposal for an AND gate starts to open the way for a very unusual type of quantum logic which is based on the flow of neutral atoms. Here, we can imagine going from AND gates to NAND gates, which are universal gates, and then by further increasing the complexity, a universal matter-wave quantum computer is accessed—at least in principle.



## B. Atomtronic SQUIDS

Quantum interference has a high importance in atomtronics, which is particularly true in the case of *atomtronic SQUIDS*. Atomtronic SQUIDS are *not* superconducting devices, but are named for their analogy with SQUIDS. They are sometimes denoted AQUIDS for Atomtronic QUAntum Interference Devices.<sup>5</sup>

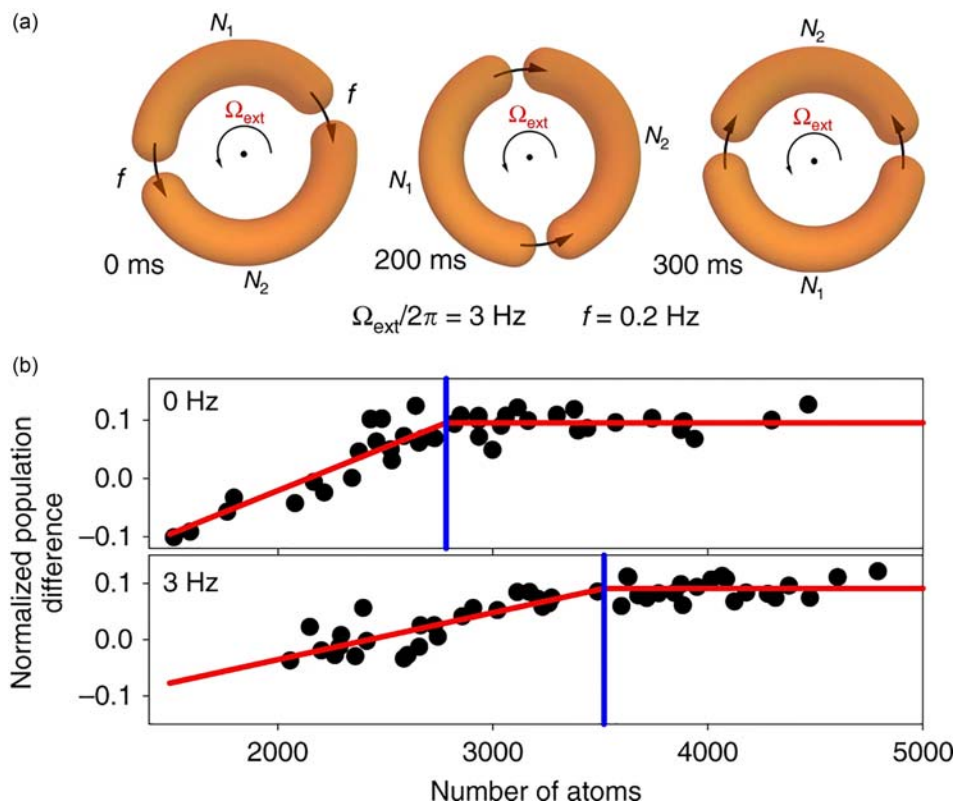
A conventional SQUID can be built from a superconducting ring with one or two “weak links,” which form Josephson junctions. At each junction, the current and the junction phase are closely related in such a way that magnetic field strengths can be determined from the oscillatory behavior of the voltage drop across the junction. The resulting device makes an excellent magnetic field sensor.<sup>507</sup>

A basic atomtronic SQUID consists of a ring waveguide for ultra-cold neutral atoms with two barriers inserted<sup>12,15</sup> [see Fig. 41(a)]. In the analogy, the ring waveguide replaces the superconducting ring, and the barriers replace the Josephson junctions. The current-phase relation for atomtronic tunnel junctions and weak links was explored in Refs. 16 and 13. Each barrier clearly affects the phase of the wavefunction and, just like the superconducting analogs, there is a critical current: in the atomtronic case, when the flow is too fast, it breaks up into vortex–antivortex pairs.<sup>67,161</sup> In a conventional SQUID, current flow in the superconducting loop is established by changing the magnetic flux through the loop. In the atomtronic SQUID, current flow in the waveguide loop is created by rotation of the system. It follows that the device’s behavior is sensitive to rotation of the SQUID (or equivalently rotation of the barriers). This principle, including quantum interference effects, has been recently demonstrated in Ref. 15. Figure

41(b) shows the transition from the AC Josephson regime (AC current through the junctions with nonzero chemical potential difference across them) to the DC Josephson regime (DC tunneling current flowing through the junctions with no chemical potential) marked with the vertical blue line as a function of the number of atoms for two different rotation rates. This transition point becomes oscillatory as a function of the rotation rate (for a fixed number of atoms) or as a function of the number of atoms (at different rotation rates as shown here). These oscillations can be used to determine the rotation rate ( $\Omega_{\text{ext}}$  in Fig. 41). Another recent work has shown that the atomtronic SQUID also exhibits hysteresis behavior analogous to the conventional SQUID.<sup>16</sup>

The recent experiments with ring traps and barriers<sup>12,15,16,67,161,391</sup> are based on optical dipole potentials. The ring trap and weak link can be created with a Laguerre–Gauss beam (with a hole) and a focused Gaussian beam.<sup>67</sup> For the ring part, potentials created by conical refraction are possible as well.<sup>145</sup> Alternatively, a very flexible approach is to use “painted” potentials,<sup>7,28</sup> where an optical dipole beam is rapidly scanned around the region of interest. As the beam is scanned, the intensity of the light is modulated so that a two-dimensional image is formed which produces a rather flexible 2D potential. Confinement in the third dimension is provided by a light sheet and the dipole potential from that. The painted potential can include a ring, and the “weak link” barriers which can be moved around the ring at will.

The atomtronic SQUID has the clear potential of being a central building block for atomtronic devices: e.g., for rotation or magnetic field sensors. However, it may also play other important roles in



**FIG. 41.** (a) We show how the “Josephson junction” barriers move in the atomtronic SQUID in order to observe a synthetic external rotation. The two barriers move at different rates  $\Omega_{\text{ext}} \pm 2\pi f$ . The number of atoms on either side of the barriers is  $N_1$  and  $N_2$ . (b) The normalized population difference  $(N_2 - N_1)/(N_2 + N_1)$  is plotted as a function of the number of atoms in each experimental run. The blue line indicates the point at which a critical atom number is reached, where the system switches from AC to DC Josephson regimes. The critical atom number varies with the “rotation” rate, being larger in the lower panel of (b). Reprinted with permission from Ryu *et al.*, Nat. Commun. 11, 3338 (2020). Copyright 2020, Author(s) licensed under a Creative Commons Attribution 4.0 License.

atomtronic circuits. It could be a component in a more complex circuit where matter wave interference is essential or allow for the storage of quantum information in the quantum states of the atomtronic SQUID.

### C. Sagnac interferometry and rotation sensing

Rotation detection devices come in many forms, including microelectromechanical systems (MEMS), hemispherical resonator gyros (HRGs), ring laser gyros (RLGs), and fiber-optic gyroscopes (FOGs). They are ubiquitous in the sense that mobile phones contain low-sensitivity gyroscopes that users can access via apps such as Phyphox. These more “traditional” sensors have been reviewed with viewpoints that are mainly academic<sup>508</sup> and industrial.<sup>509</sup>

Atoms can also be used as gyroscopes, for example, via nuclear spins in comagnetometers.<sup>510</sup> Alternatively, the demonstration of atom-optical manifestations of the Sagnac effect dated in 1991,<sup>511</sup> and already in 1997, short-term sensitivities of  $2 \times 10^{-8}$  (rad/s)/ $\sqrt{\text{Hz}}$  were demonstrated in an atom-based Sagnac interferometer,<sup>50,512</sup> since then there have been many technical advances in such “free space” interferometers,<sup>513</sup> commonly based on a Mach–Zehnder-type configuration.

From an atomtronic standpoint, we wish to consider “closed-path” guided configurations, which have been used for cold molecules,<sup>514</sup> cold atoms,<sup>101,155</sup> and Bose–Einstein condensates<sup>103,515</sup> (BECs) since the early 2000s and have been recently considered for chip-scale development.<sup>108,173</sup> Theoretical and experimental guiding geometries include optical dipole, magnetic and Stark confinement using constant, time-averaged, inductive or dressed potentials.<sup>6,7,9,28,32,102,107,142,143,158,390,406,516–535</sup> Highly supersonic superfluid flow is also possible;<sup>103,117,128</sup> however, while the concept of closed-path cold atom configurations has long held traction, the sensitivity has yet to reach the high levels of free-space cold atom gyros.<sup>536</sup> There has been some important recent experimental progress on this front,<sup>537,538</sup> i.e., all the necessary experimental tools appear, in principle, to be present.

This means we consider Bose-condensed atoms held within an appropriate (e.g., toroidal, as used in atomtronic SQUIDs) trapping potential  $V$ . This is necessarily assumed to be in a rotating frame defined by the angular velocity vector  $\Omega$ , manifesting as an additional term  $i\hbar\Omega \cdot (\mathbf{x} \times \nabla)$  in either the single-particle Schrödinger equation or the Gross–Pitaevskii equation (GPE) when describing mean-field dynamics of Bose-condensed atoms. A coordinate system where  $\Omega$  points along the  $z$  axis (dynamics viewed from within a frame rotating anticlockwise around the  $z$  axis with angular frequency  $\Omega$ ) simplifies this term to  $i\hbar\Omega(x\partial_y - y\partial_x)$ . A good general starting point for describing the dynamics within a variety of such systems is the following system of GPEs,

$$i\hbar \frac{\partial}{\partial t} \Psi_j = -\frac{\hbar^2}{2m} \nabla^2 \Psi_j + i\hbar\Omega \left( x \frac{\partial}{\partial y} \Psi_j - y \frac{\partial}{\partial x} \Psi_j \right) + V_j \Psi_j + \sum_j g_{jj} |\Psi_j|^2 \Psi_j. \quad (67)$$

The  $j$  index labels different internal atomic states, the  $V_j$  incorporate energy differences between the internal states and any internal state dependences of the trapping potential, and the  $g_{jj}$  quantify the strengths of  $s$ -wave scattering terms (ignoring the possibility of internal-state-changing collisions).

We first consider a tight toroidal trapping potential, reducing our treatment to a radius  $R$  one-dimensional ring geometry, leaving only the polar angle  $\phi$  free, reduce to a single internal state, and neglect all interactions. The GPE then becomes

$$i\hbar \frac{\partial}{\partial t} \psi(\phi) = \left( -\frac{\hbar^2}{2mR^2} \frac{\partial^2}{\partial \phi^2} + i\hbar\Omega \frac{\partial}{\partial \phi} \right) \psi(\phi). \quad (68)$$

Note that the interactions are genuinely insignificant if the gas is very dilute, or if interactions are tuned away using an appropriate Feshbach resonance. The evolution of an initial, localized matter wave split into an equal superposition with opposite velocity splitting products can, at the simplest level, be considered without explicit mention of the initial wave packet. At this level, an initial state  $\psi(0)$  evolves as  $\psi(t)$ :

$$\begin{aligned} \psi(0) &= (e^{i\ell\phi} + e^{-i\ell\phi})/\sqrt{4\pi} = \cos(\ell\phi)/\sqrt{\pi}, \\ \psi(t) &= e^{-i\hbar\ell^2 t/2mR^2} \cos(\ell[\phi + \Omega t])/\sqrt{\pi}, \end{aligned}$$

yielding intensity fringes  $\propto (2/\pi)\{\cos(2\ell[\phi + \Omega t]) + 1\}$  multiplied by the number of atoms whenever the matter wave splitting products overlap in space. For a matter wave initially centered at  $\phi = 0$  enclosed in a ring with area  $A = \pi R^2$ , this occurs at  $\phi = \pi$  or  $0$ , when  $t = (1 \text{ or } 1)Am/\hbar\ell$ , yielding a phase shift of  $\Delta\phi = \Omega t = (2 \text{ or } 4)A\Omega m/\hbar$ , respectively.

Using the speed  $v = \hbar\ell/Rm$  and wavelength  $\lambda = 2\pi R/\ell$  of the propagating atoms, we express the latter phase shift as  $\Delta\phi = 8\pi A\Omega/\lambda v$ , the same form as the phase shift accumulated by an optical Sagnac interferometer, with  $\lambda$  and  $v$  replaced by the wavelength and speed of light, respectively. This highlights the promise of atom interferometry, in that  $\lambda v$  can be made much smaller than its optical equivalent. Also, note that if the initial state is literally  $\psi(0) = \cos(\ell\phi)/\sqrt{\pi}$ , i.e., there is no localizing “envelope” to the wave packet, the accrued fringe shift can be observed at any time, effectively increasing the enclosed area of the interferometer. This highlights an important feature in that as the speed of light is a constant, it is necessary to increase  $A$  in order to increase the interrogation time; with cold atoms, this is not the case.

With typical repulsive interactions (positive  $g$ ), such standing wave fringes will rapidly disperse; however, in a two-component system with very similar scattering lengths ( $\Rightarrow g_{11} \approx g_{12} \approx g_{22}$ , as can be achieved in <sup>87</sup>Rb),<sup>539</sup> producing an equal superposition initial state such that  $\psi_1(0) = \cos(\ell\phi)/\sqrt{2\pi}$ ,  $\psi_2(0) = \sin(\ell\phi)/\sqrt{2\pi}$ , the two components stabilize each other by making the total mean field potential  $g(|\psi_1|^2 + |\psi_2|^2)$  essentially flat (hence, no gradients and no dispersive forces). Alternatively, initializing the system such that  $\psi_1(0) = e^{i\ell\phi}/\sqrt{4\pi}$ ,  $\psi_2(0) = e^{-i\ell\phi}/\sqrt{4\pi}$ , followed by an evolution time  $T/2$ , a  $\pi$  pulse swapping the internal states, and a second evolution time  $T/2$ , produces

$$\begin{aligned} \psi_1(T) &= e^{-i\varphi T} e^{-i\ell(\phi + \Omega T)}/\sqrt{4\pi}, \\ \psi_2(T) &= e^{-i\varphi T} e^{i\ell(\phi + \Omega T)}/\sqrt{4\pi} \end{aligned}$$

( $\varphi$  is a global phase depending on  $\ell^2$  and the values of  $g_{11}$ ,  $g_{12}$ , and  $g_{22}$ ). Repeating the initializing process produces

$$\begin{aligned} \psi_1(T) &= -ie^{-i\varphi T} \sin(\ell[\phi + \Omega T])/\sqrt{2\pi}, \\ \psi_2(T) &= e^{-i\varphi T} \cos(\ell[\phi + \Omega T])/\sqrt{2\pi}. \end{aligned}$$

The value of  $\Omega$  can then be inferred from population measurements:  $N_1 = N[1 + \cos(2\ell\Omega T)]/2$ ,  $N_2 = N[1 - \cos(2\ell\Omega T)]/2$ , where  $N$  is the total particle number. The  $\pi$  swap pulse at  $T/2$  is carried out to counteract accumulation of relative phase due to differences in internal state energy and values of  $g_{11}$  and  $g_{22}$ ; if  $g_{11} \approx g_{22}$  and the energy gap between the internal states is well known, this can be neglected, and a detailed experimental proposal based around  $^{87}\text{Rb}$  and magnetic vortex pumping has been determined by Helm *et al.*<sup>540</sup> This alternative has the advantage of there being no mean field gradients even when the scattering lengths are quite different and turns a measurement of interference fringes into a measurement of relative population. Note, however, that optimum sensitivity of such a measurement is when the slope of the response curve is maximal, e.g., when  $2\ell\Omega T \approx \pi/2$ ;<sup>540,544</sup> it may, therefore, be advisable to add a controlled relative phase in an experimental realization.

A quite different approach, again in near 1D, is to have a single component condensate with attractive interactions, using, e.g.,  $^{85}\text{Rb}$  or  $^7\text{Li}$ , again ideally in a ring geometry, as illustrated in Fig. 42.<sup>542</sup> In this case, the GPE exhibits soliton solutions: stable, nondispersive, localized wave packets which are robust to collisions, behaving something like classical particles. Given a sufficiently sharp barrier—ideally a  $\delta$ -function, more realistically a Gaussian barrier with width significantly smaller than the soliton's characteristic length<sup>543,552</sup> formed, e.g., by a focused off-resonant sheet of light<sup>545</sup>—an initial soliton can be split into two halves propagating with equal and opposite velocity if its incoming velocity is correctly calibrated to the barrier size. In an essentially similar way, the splitting products can accumulate a relative Sagnac phase, which could, in principle, be visualized through spatial interference fringes.<sup>546</sup> The solitons' small size can make this a less

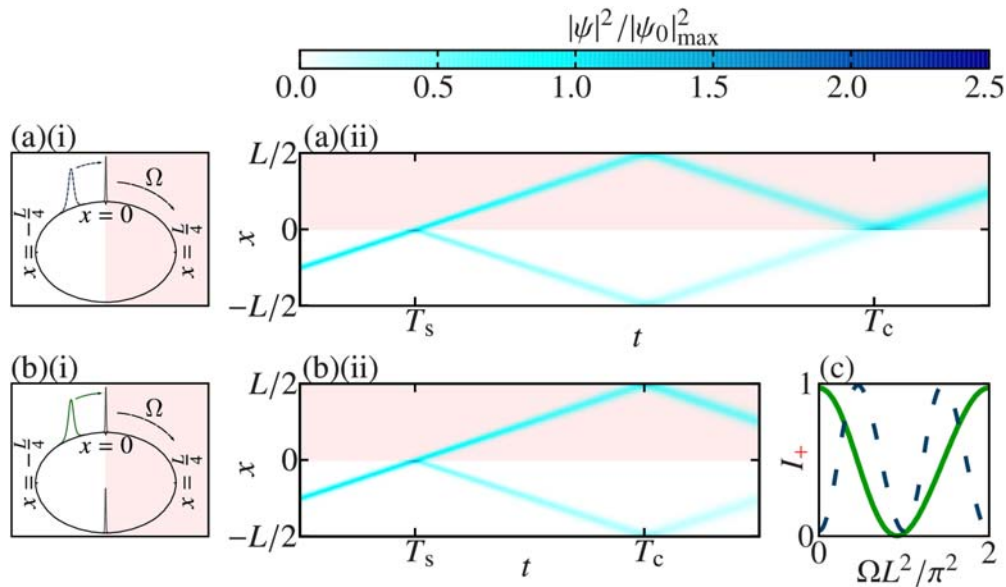
suitable approach, however, than recombining them again on a barrier, where the relative phase manifests through the relative sizes of the wave packets emerging on either side of the barrier as the result of this second collision, i.e., again as relative population measurement, with an essentially similar (ideal) dependency on  $\Omega$  as that outlined above. This second barrier interaction can take place either at a second barrier exactly opposite to the first,<sup>547</sup> or due to the fact that solitons are robust to collisions and, therefore, in some sense they should “pass through” one another back at the same barrier at which the initial splitting took place, following both splitting products having completely circumnavigated the ring.<sup>542</sup>

The roles of quantum noise and interaction for rotation sensing with bright solitons in the quantum regime were studied in Ref. 548. It was found that interaction and noise should be carefully considered in order the performances of the system are not spoiled. In Ref. 549, the scattering properties of a quantum matter wave soliton splitting in a barrier were studied. In addition, the GPE analysis is of limited accuracy for the quantitative analysis of the sensitivity of atom interferometry in the presence of interaction. For other features of bright soliton interferometers, please see Sec. XIV.

Finally, we note that everything we have described is in a sense “classical,” in the sense of a classical field description of the BEC being completely adequate and that, more explicitly, quantum elaborations have been proposed, exploiting spin squeezing<sup>550</sup> or ideas from quantum information.<sup>551</sup>

#### D. Magnetometry

The development of compact highly sensitive magnetometers with high spatial resolution is one of the current challenges of



**FIG. 42.** An incoming soliton splits at time  $T_s$  on a barrier into two solitons of equal amplitude and opposite velocity. At a time  $T_c$ , the solitons recombine either at the same barrier (a), or a second barrier (b) antipodal to the first (the example value of  $\Omega$  is the same in both cases). The resulting phase difference is read out via the population difference in the final output products within the positive (shaded) and negative domains. (c) Final population in the positive domain  $I_+$  as a function of  $\Omega$ . The sensitivity of the single barrier case (dashed line) is twice that of the double barrier case (solid line) because the interrogation time  $T_c - T_s$  is doubled. Reprinted with permission from Helm *et al.*, Phys. Rev. Lett. **114**, 134101 (2015). Copyright 2015, American Physical Society.



Atomtronics. The capability of measuring very weak magnetic fields with high precision and accuracy is at the basis of numerous applications, including bio-magnetism, geology, data storage, and archaeology.<sup>552</sup> Different approaches have been followed in the previous years to reach this goal, mainly using superconducting quantum interference devices (SQUIDS), nitrogen-vacancy (NV) diamond magnetometers, and atomic magnetometers. Currently, SQUIDS and spin-exchange relaxation-free magnetometers reach sensitivities at the  $\text{fT}/\sqrt{\text{Hz}}$  level and below,<sup>553</sup> whereas NV-magnetometers allow for  $\text{pT}/\sqrt{\text{Hz}}$  sensitivities.<sup>554</sup> A valuable pictorial log-log plot of experimental device scale vs magnetic sensitivity for a wide variety of magnetometer sensor technologies can be found in Fig. 2 of a recent review.<sup>555</sup>

Atomic magnetometers can be classified depending on whether the magnetic field drives the internal or the external degrees of freedom of the atoms. The former are typically based on the measurement of the Larmor spin precession of optically pumped atoms either using thermal clouds or BECs. In the case of thermal clouds, double-resonance optically pumped magnetometers are an attractive instrument for unshielded magnetic-field measurements due to their wide dynamic range and high sensitivity.<sup>556</sup> In the BEC case, the use of stimulated Raman transitions has been reported<sup>557</sup> as well as the separate probe of the different internal states of a spinor BEC after free fall<sup>558</sup> or the measure of the Larmor precession in a spinor BEC.<sup>559–564</sup> Also, note that a two-component BEC has been also investigated for magnetometry.<sup>565</sup>

An alternative approach to atomic magnetometry is based on encoding the magnetic field information in the spatial density profile of matter waves. Some examples of this approach are those based on detecting density fluctuations in a BEC due to the magnetic induced deformation of the trapping potential.<sup>237,566,567</sup> Recently, a different scenario has been explored and a quantum device for measuring two-body interactions, scalar magnetic fields and rotations based on a BEC in a ring trap has been proposed.<sup>568</sup> To this aim, the BEC is prepared in an imbalanced superposition of the two counter-propagating orbital angular momentum (OAM)  $l = 1$  modes and due to quantum interference, a line of minimal atomic density appears. In the presence of nonlinear interactions, this nodal line shows a solitonlike rotating motion (Fig. 43). An analytical expression relating the angular frequency of the rotation of the minimal density line,  $\Omega_m$ , to the strength of the nonlinear atom–atom interactions and the difference between the populations of the counter-propagating modes is derived,

$$\Omega_m = \frac{Un_{1\pm}}{2\left(1 + \frac{U}{\Delta}\right)}, \quad (69)$$

where  $n_{1\pm}$  is the population imbalance between the  $l = \pm 1$  modes,  $U$  is a nonlinear parameter proportional to the scattering length, and  $\Delta$  is the chemical potential difference between the  $l = 3$  and  $l = 1$  modes, see Ref. 568. This expression constitutes the basis to use the physical system under consideration as a quantum sensing device by measuring the rotation frequency of the minimum density line by direct imaging, in real time, the spatial density distribution of the BEC. In fact, a full experimental protocol based on direct fluorescence imaging of the BEC that allows one to measure all the quantities involved in the analytical model is proposed.

Let us assume that the lifetime of the BEC is  $\tau$ . Then, the condition  $\Omega_m \omega \geq 1/\tau$  being  $\omega$ ; the ring trapping frequency must be fulfilled to be able to observe the rotation of the minimum density line for the

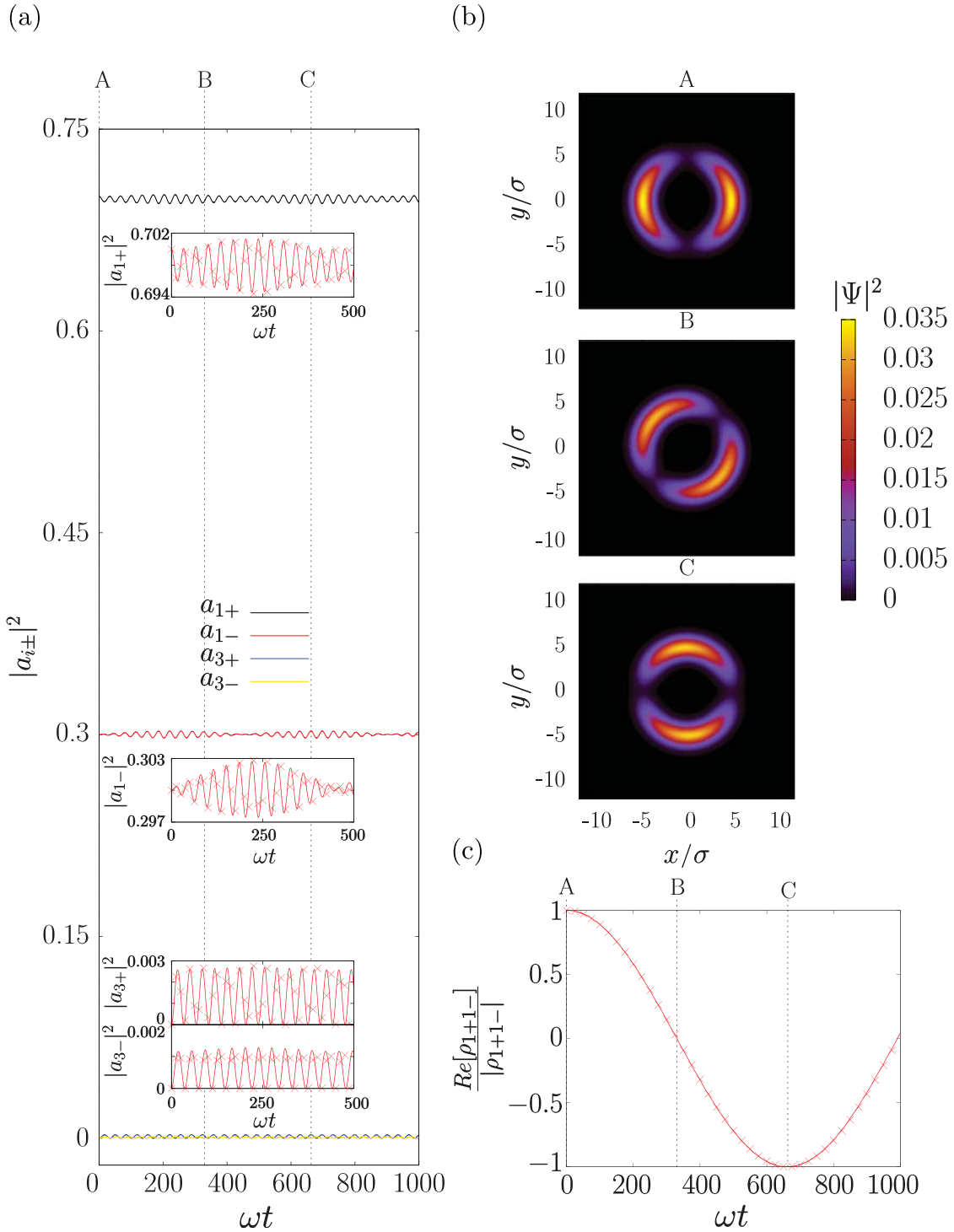
time the experiment lasts. The upper limit of observable values of  $\Omega$  is imposed by the regime of validity of the model, i.e.,  $\Omega_m < 0.025$  to avoid the excitation of states with OAM higher than 1. A magnetic field produces, in general, a variation of the scattering length, which does not depend on the magnetic field orientation. Therefore, the presence of an external magnetic field will induce a variation of  $\Omega_m$  and the system could be used as a scalar magnetometer by relating changes on the frequency of rotation of the minimal line to variations of the modulus of the magnetic field. The sensitivity in the measurement of magnetic field variations increases with the number of condensed particles but keeping the scattering length small and having a strong dependence of the scattering length on the magnetic field modulus. Thus, close to a Feshbach resonance, these requirements could be met. However, close to a Feshbach resonance, the three-body losses may limit the lifetime of the BEC making difficult the measurement. Nevertheless, some atomic species such as  $^{85}\text{Rb}$ ,  $^{133}\text{Cs}$ ,  $^{39}\text{K}$ , or  $^7\text{Li}$  have been reported to form BECs that are stable across Feshbach resonances with lifetimes on the order of a few seconds, so they could be potential candidates for using the system as a magnetometer. Taking into account that the trapping frequency  $\omega$ , is typically of the order of a few hundreds of Hz for ring-shaped traps, and considering typical values of  $\Omega_m \omega \sim 1$  Hz compatibles with typical times for the experiment of around  $\tau \sim 1$  s, the minimum density line would perform some complete round trips. Assuming that one could resolve angular differences on the order of  $\sim 0.1$  rad, variations in the rotation frequency on the order of  $10^{-2}$  Hz could be measured. Thus, for the parameter values reported in Ref. 569, in principle, this magnetometer would allow one to measure changes in the magnetic field on the order of a few pT at a bandwidth of 1 Hz.

## E. Concluding remarks and outlook

In this section, we have presented some examples for engineering of atomtronic devices. We have discussed recent advances in the development of basic components for atomtronics such as batteries, diodes, and transistors. However, atomtronics applications are expected to go beyond analogs of electronic circuits with atoms by making use of the specific quantum properties of ultracold atomic matter. In this context, we have shown that by taking advantage of quantum coherence and interactions between ultracold atoms, it is possible to design atomtronic SQUIDS, matter-wave interferometers, as well as rotation and magnetic sensors with extremely high accuracy and resolution. All these applications open the door to the future development of completely new types of quantum devices, which might be integrated into complex and large-scale atomtronic circuits.

## ACKNOWLEDGMENTS

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**FIG. 43.** (a) Time evolution of the population of the states involved in the dynamics. (b) Snapshots of the density profile for different instants of the dynamical evolution. (c) Time evolution of the real part of the coherence between the  $|1, +\rangle$  and  $|1, -\rangle$  states. The points correspond to the numerical simulation of the GPE, while the continuous lines are obtained by solving the FSM equations. The considered parameter values are  $R=5$ ,  $g_{2d}=1$ , for which  $U=0.0128$ ,  $\mu_1=0.529$  and  $\mu_3=0.699$ ,  $a_{1+}(0)=\sqrt{p_{1+}(0)}=\sqrt{0.7}$  and  $a_{1-}(0)=\sqrt{p_{1-}(0)}=\sqrt{0.3}$ . Reprinted with permission from Pelegrí *et al.*, New J. Phys. **20**, 103001 (2018). Copyright 2018, Author(s) licensed under a Creative Commons Attribution 3.0 License.

## X. TWO LEVEL QUANTUM DYNAMICS IN RING-SHAPED CONDENSATES AND MACROSCOPIC QUANTUM COHERENCE

**D. Aghamalyan, M. Boshier, R. Dumke, T. Haug, A. Minguzzi, L.-C. Kwek, L. Amico**

A qubit is a two state quantum system that can be coherently manipulated, coupled to its neighbors, and measured. Several qubit physical implementations have been proposed in the last decade, all of them presenting specific virtues and bottlenecks at different levels.<sup>570–575</sup> In neutral cold atoms proposals, the qubit is encoded into well isolated internal atomic states. This allows long coherence times, precise state readout, and, in principle, scalable quantum registers. However, an individual qubit (atom) addressing is a delicate point.<sup>576,577</sup> Qubits based on Josephson junctions allow fast gate operations and make use of the precision reached by lithography techniques.<sup>578</sup> The decoherence, however, is fast in these systems, and it is experimentally challenging to reduce it. For charged qubits, the main problem arises from dephasing due to background charges in the substrate; flux qubits are insensitive to the latter decoherence source but are influenced by magnetic flux fluctuations due to impaired spins proximal to the device.<sup>579</sup>

Here, we aim at combining the advantages of cold atom and Josephson junction based implementations. The basic idea is to use the persistent currents flowing through ring shaped optical lattices<sup>13,14,401,579–583</sup> to realize a cold atom analog of the superconducting flux qubit (see Refs. 165, 579, and 584–587 for the different schemes that can be applied to induce persistent currents). A barrier potential painted along the ring gives rise to a weak link, acting as a source of back-scattering for the propagating condensate, thus creating an interference state with the forward scattered current. This gives rise to an atomic condensate counterpart of the celebrated rf-SQUID—a superconducting ring interrupted by a Josephson junction,<sup>445,570</sup> namely, an Atomtronic Quantum Interference Device (AQUID). Due to the promising combination of advantages characterizing Josephson junctions and cold atoms, the AQUID is now an object of intense investigation.<sup>16,391</sup> The first experimental realizations have been done by means of a Bose–Einstein condensate free to move along a toroidal potential, except through a small spatial region, where a very focused blue-detuned laser creates weak links, namely, an effective potential constriction.<sup>12,67,161</sup> By adapting the logic applied in the context of solid state Josephson junctions<sup>570,588</sup> to a specific cold atoms setup, a cold atom version of the SQUID can be created. On the theoretical side, it has been demonstrated that the two currents flowing in the AQUID can, indeed, define an effective two-level system, that is, the cold-atom analog of flux qubits.<sup>13,14,579,580,589</sup> The system is assumed to be driven by an effective flux piercing the ring lattice. The potential constriction breaks the Galilean invariance and splits the qubit levels, that otherwise would be perfectly degenerate at half-flux quantum. By a combination of analytic and numerical techniques, one can demonstrate that the system can sustain a two-level effective dynamics.<sup>13,14,579,580</sup> We also review a physical system consisting of a Bose–Einstein condensate confined to a ring shaped lattice potential interrupted by three weak links.<sup>14</sup> By employing path integral techniques, we explore the effective quantum dynamics of the system in a pure quantum phase dynamics regime. By a combination of analytic and numerical techniques, it was demonstrated that the

system can sustain a two-level effective dynamics giving other realization of the atomtronic qubit.

After outlining theoretical framework, which leads to obtaining a single qubit, we go further by showing how single qubit and two-qubit gates can be implemented by using an effective action approach.<sup>401,580</sup> In order to achieve two-qubit gates, we allow a nonvanishing hopping term between the different rings.

We also review the experimental realization of ring lattices with one and three weak links performed at Nanyang Technological University in the experimental group of Dumke in Singapore. Indeed using a spatial light modulator, they have experimentally realized<sup>580</sup> both the single ring lattice with a weak link and scaled ring-lattice potentials that could host, in principle,  $n \geq 10$  ring-qubits, arranged in a stack configuration, along the laser beam propagation axis. Trapping potential of a ring-shaped optical lattice with three weak links a  $\sim 20 \mu\text{m}$  diameter using a spatial light modulator has been reported in Ref. 13.

The current and type of state inside these atomic qubits can be read out via time-of-flight measurements.<sup>590</sup> When the ring is interfered with a reference condensate, a spiral pattern appears in the time of flight, which indicates the magnitude and direction of the current. For low resolution images, these spirals can be read out from the density–density correlation images. Furthermore, the type of superposition state can be measured from the noise in the time-of-flight images.<sup>590</sup>

First progress toward an experimental realization has been made recently. In a recent experiment, interference of persistent currents of AQUIDs has been demonstrated at the Los Alamos National Laboratory by Ryu *et al.*<sup>15</sup> By inducing a bias current in a rotating atomic ring interrupted by two weak links, the interference between the Josephson current with the current from the rotation creates a oscillation in the critical current with applied flux. This oscillation is measured experimentally in the transition from the DC to the AC Josephson effect. This experiment has been performed within a dilute Bose–Einstein condensate that is well described within a mean-field description and thus entanglement of currents, which is a key ingredient for the atomic qubit, has not been demonstrated. Nonetheless, it is a major step toward the implementation of the atomic qubit.

### A. The Atomtronic quantum interference device: AQUID

We start by considering analytical models for the confined one-dimensional many-body systems and use them to demonstrate an emerging effective two level dynamics of the system. Let us start by considering  $N$  interacting bosons at zero temperature, loaded into a 1D ring-shaped optical lattice of  $M$  sites. The discrete rotational symmetry of the lattice ring is broken by the presence of a localized potential in one lattice site (later we also consider case of three weak links), which gives rise to a weak link. The relevant physics of the system is captured by the Bose–Hubbard model. The Hamiltonian reads

$$\mathcal{H}_{\text{BH}} = \sum_{i=1}^M \left[ \frac{U}{2} n_i(n_i - 1) + \Lambda_i n_i - J_i \left( e^{-i2\pi\Omega/M} a_{i+1}^\dagger a_i + \text{h.c.} \right) \right], \quad (70)$$

where  $a_i$  ( $a_i^\dagger$ ) are bosonic annihilation (creation) operators on the  $i$ th site of a ring with length  $M$  and  $n_i = a_i^\dagger a_i$  is the corresponding number operator. Periodic boundaries are imposed, meaning that



$a_M \equiv a_0$ . The parameter  $U$  takes into account the finite scattering length for the atomic two-body collisions on the same site:  $U = 4\pi\hbar^2 a_0 \int dx |w(x)|^4 / m$ , with  $w(x)$  being the Wannier functions of the lattice,  $m$  the mass of atoms, and  $a_0$  the scattering length. To break the translational symmetry, there are two possible ways: Either, the hopping parameters are all equal  $J_i = J$  except in one weak-link hopping  $i_0$  where  $J_{i_0} = J'$ . The other alternative, which we choose in this review, is to place a potential barrier at a single site  $\Lambda_i = \Lambda$ , and at all other sites, the potential is set to zero with  $J_i = J, \forall i$ . The two options show qualitatively the same physics.<sup>14</sup> The ring is pierced by an artificial (dimensionless) magnetic flux  $\Omega$ , which can be experimentally induced for neutral atoms as a Coriolis flux by rotating the lattice at constant velocity<sup>12,67,161</sup> or as a synthetic gauge flux by imparting a geometric phase directly to the atoms via suitably designed laser fields.<sup>586,591</sup> The presence of the flux  $\Omega$  in Eq. (70) has been taken into account through the Peierls substitution:  $J_i \rightarrow e^{-i2\pi\Omega/M} J_i$ . The Hamiltonian [Eq. (75)] is manifestly periodic in  $\Omega$  with period 1. In the absence of the weak-link, the system is also rotationally invariant and, therefore, the particle-particle interaction energy does not depend on  $\Omega$ . The many-body ground-state energy, as a function of  $\Omega$ , is, therefore, given by a set of parabolas intersecting at the frustration points  $\Omega_n = (n + \frac{1}{2})$ .<sup>476,592</sup> The presence of the weak-link breaks the axial rotational symmetry and couples different angular momenta states, thus lifting the degeneracy at  $\Omega_n$ . This feature sets the qubit operating point.<sup>14,580</sup>

It is worth noting that the interaction  $U$  and the weak-link strength induce competing physical effects: the weak-link sets an healing length in the density as a further spatial scale; the interaction tends to smooth out the healing length effect. As a result, strong interactions tends to renormalize the weak link energy scale.<sup>14,406,581</sup>

In the limit of a large number of bosons in each well  $\bar{n} = N/M$ ,  $a_i \sim \sqrt{\bar{n}} e^{i\phi_i}$  and the Bose-Hubbard hamiltonian (BHH), Eq. (70) can be mapped to the quantum-phase model employed to describe Josephson junction arrays:<sup>593,594</sup>

$$\mathcal{H} = \sum_{i=1}^M \left[ \frac{U}{2} \mathbf{n}_i^2 - J_i \cos(\phi_{i+1} - \phi_i - \Omega) \right], \quad (71)$$

where  $[\mathbf{n}_i, \phi_i] = i\hbar\delta_{ij}$  are canonically conjugate number-phase variables and  $J_i \sim \bar{n} t_i$  are the Josephson tunneling amplitudes.

### 1. The rf-AQUID qubit

In this case, a single weak link occurs along the ring lattice  $t'' = t$ . The presence of the weak link induces a slow/fast separation of the effective (imaginary time) dynamics: the dynamical variables relative to the weak link are slow compared to the “bulk” ones, playing the role of an effective bath (nonetheless, we assume that the ring system is perfectly isolated from the environment). Applying the harmonic approximation to the fast dynamics and integrating it out, the effective dynamics of the AQUID is governed by (see for detailed derivation appendix material of Ref. 580),

$$\mathcal{H}_{\text{eff}} = \mathcal{H}_{\text{sys}} + \mathcal{H}_{\text{bath}} + \mathcal{H}_{\text{sys-bath}}. \quad (72)$$

The slow dynamics is controlled by

$$\mathcal{H}_{\text{sys}} = U \mathbf{n}^2 + E_L \varphi^2 - E_J \cos(\theta - \Omega), \quad (73)$$

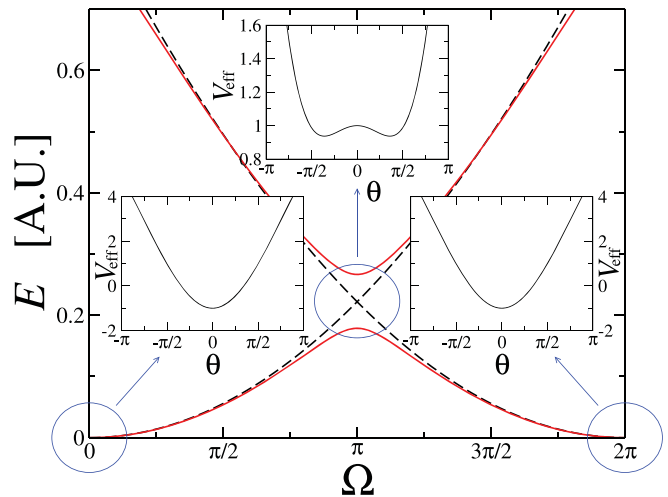
where  $\theta$  is the phase slip across the weak link with  $E_L = J/M$  and  $E_J = J'$ . For  $\delta \equiv E_J/E_L \geq 1$ ,  $\mathcal{H}_{\text{sys}}$  describes a particle in a double well

potential with the two-minima-well (see Fig. 44) separated from the other features of the potential. The two parameters,  $U$  and  $t'/t$ , allow control of the two level system. The two local minima of the double well are degenerate for  $\Omega = \pi$ . The minima correspond to the clockwise and anticlockwise currents in the AQUID. The presence of a finite barrier,  $\Lambda > 0$ , breaks the axial rotational symmetry and couples different angular momenta, thus lifting the degeneracy at the frustration points by an amount  $\Delta E$ , see Fig. 44. Provided other excitations are energetically far enough from the two competing ground-states, this will identify the two-level system defining the desired qubit and its working point. Because of the quantum tunneling between the two minima of the double well, the two states of the system (qubit) are formed by symmetric and antisymmetric combinations of the two circulating current states.

The WKB level splitting is (see for detailed derivation in Appendix C.3 of Ref. 595)

$$\Delta \simeq \frac{2\sqrt{UE_J}}{\pi} \sqrt{\left(1 - \frac{1}{\delta}\right)} e^{-12\sqrt{E_J/U}(1-1/\delta)^{3/2}}. \quad (74)$$

From this formula, we can see that the limit of a weak barrier and intermediate to strong interactions form the most favorable regime to obtain a finite gap between the two energy levels of the double level potential as depicted on Fig. 44. Incidentally, we comment that the bath Hamiltonian in Eqs. (72) and (73) is similar to the one describing the dissipative dynamics of a single Josephson junction in the framework of the Caldeira-Leggett model.<sup>476</sup> As long as the ring has a finite size, however, there are a finite number of discrete modes and no real



**FIG. 44.** Main panel: sketch of the qubit energy splitting, due to the barrier  $\Lambda$ , for the two lowest-lying energy states in the many-body spectrum of model (70). Black dashed lines denote the ground-state energy in the absence of the barrier, as a function of the flux  $\Omega$ . Switching on the barrier opens a gap at the frustration point  $\Omega = \pi$  (continuous red lines). The three insets show the qualitative form of the effective potential at  $\Omega = 0, \pi, 2\pi$ . Note the characteristic double-well shape forming at  $\Omega = \pi$ . The qubit, or effective two-level system, corresponds to the two lowest energy levels of this potential. In this figure the energies are plotted in arbitrary units. Reprinted with permission from Aghamalyan *et al.*, New J. Phys. 17, 045023 (2015). Copyright 2015, Author(s) licensed under a Creative Commons Attribution 3.0 License.

dissipation occurs.<sup>596</sup> In the limit  $N \rightarrow \infty$ , a proper Caldeira–Leggett model is recovered. In agreement to the arguments reported above, the qubit dynamics encoded in the AQUID is less and less addressable by increasing the size of the ring.<sup>14,580</sup>

## 2. Atomtronic flux-qubit: Ring lattice interrupted with three weak links

Here, we consider  $N$  Bosons in an  $M$  site ring described by the Bose–Hubbard model. The Hamiltonian reads

$$\mathcal{H}_{\text{BHH}} = \sum_{i=1}^M \left[ \frac{U}{2} n_i(n_i - 1) - t_i \left( e^{i\Omega} a_{i+1}^\dagger a_i + \text{h.c.} \right) \right]. \quad (75)$$

where  $a_i$  ( $a_i^\dagger$ ) are bosonic annihilation (creation) operators on the  $i$ th site and  $n_i = a_i^\dagger a_i$  is the corresponding number operator. Periodic boundaries are imposed, meaning that  $a_{M+1} \equiv a_1$ . The parameter  $U$  takes into account the finite scattering length for the atomic two-body collisions on the same site. The hopping parameters are constant  $t_j = t$  except in the three weak-links lattice sites  $i_0, i_1, i_2$ , where they are  $t_{i_0} = t'$ ,  $t_{i_1} = t_2 = t''$ . The ring is pierced by an artificial (dimensionless) magnetic flux  $\Omega$ , which can be experimentally induced for neutral atoms as a Coriolis flux by rotating the lattice at constant velocity,<sup>161,597</sup> or as a synthetic gauge flux by imparting a geometric phase directly to the atoms via suitably designed laser fields.<sup>591,598,599</sup> The presence of the flux  $\Omega$  in [Eq. (75)] has been taken into account through the Peierls substitution:  $t_i \rightarrow e^{-i\Omega} t_i$ . The Hamiltonian [Eq. (75)] is manifestly periodic in  $\Omega$  with period  $2\pi$ ; in addition, it enjoys the symmetry  $\Omega \leftrightarrow -\Omega$ . The presence of the weak-link breaks the axial rotational symmetry and couples different angular momenta states, thus lifting the degeneracy at  $\Omega_n$ . This feature sets the qubit operating point.<sup>14,580</sup>

Here, again in the limit of a large number of bosons in each well  $\bar{n} = N/M$ ,  $a_i \sim \sqrt{\bar{n}} e^{i\phi_i}$ , and the Bose–Hubbard hamiltonian (BHH) [Eq. (75)] can be mapped to the quantum-phase model and equivalent Hamiltonian is given by Eq. (71).

The effective action for the quantum phase model reads (see Ref. 13 for details),

$$S_{\text{eff}} = \sum_{\alpha=0,1,2} \int_0^\beta d\tau \left[ \frac{1}{U} \dot{\theta}_\alpha^2 + V(\theta_\alpha) \right] \quad (76)$$

$$- \frac{J}{U} \int d\tau d\tau' \theta_\alpha(\tau) G_\alpha(\tau - \tau') \theta_\alpha(\tau'), \quad (77)$$

where

$$V(\theta_\alpha) \doteq J c_\alpha \theta_\alpha^2 - \frac{J'}{3} \cos(\theta_1 - \theta_2 - \Omega) - \frac{J''}{3} (\cos \theta_1 + \cos \theta_2), \quad (78)$$

with  $c_\alpha = \frac{1}{2} (1 - UJ \sum_{k=1}^{M-4} \frac{\zeta_\alpha k^2}{\omega_k^2})$ , where  $\theta_\alpha \doteq \phi_{i_\alpha+1} - \phi_{i_\alpha}$ . We assume that the weak links are sufficiently spaced to make the nearest neighbor phase differences in between them (fast variables) small. This implies that substantial phase slips occur at the weak links with the constraint  $\theta_0 + \theta_1 - \theta_2 = 0 \bmod(2\pi)$ . The interaction between the fast and the slow modes is described by the kernel,

$$G_\alpha(\tau) = \sum_{l=0}^{\infty} \sum_{k=1}^{M-4} \frac{\omega_l^2 \zeta_\alpha k^2}{\omega_k^2 + \omega_l^2} e^{i\omega_l \tau}. \quad (79)$$

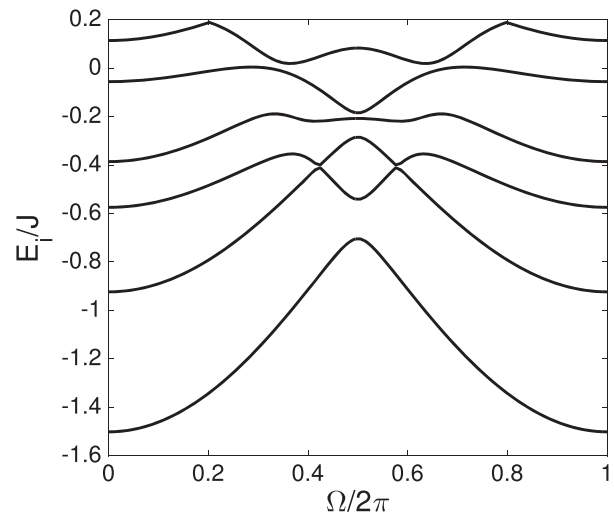
We observe that  $V(\theta_\alpha)$  defines the effective dynamics of the superconducting Josephson junctions flux qubits,<sup>588,600</sup> but perturbed by the  $\theta^2$  terms; by numerical inspection, we see that the corresponding coefficients are small in units of  $J$  and decrease by increasing  $M$ . Moreover, in Fig. 45, we introduce the numerical result for the spectrum of the quantum particle that moves in the potential given by Eq. (79) under the additional assumption that  $\theta^2$  terms do not contribute. From this figure, we clearly see that near the frustration point  $\Omega = \pi$  two lowest energy levels are well separated from each other and from higher excitations, which means that effective dynamics of the system defines a qubit. It is important to point out that quantum phase model is applicable in the limit of the high filling; however, the results for the effective-two level description were demonstrated to hold in the limit of low filling by applying an exact diagonalization method for the Bose–Hubbard model as it has been demonstrated in Refs. 14 and 15

## B. Demonstration of the one qubit and two qubit unitary gates

The aim of this section is to show how the effective phase dynamics of optical ring-lattices with impurities serves to the construction of one- and two-qubit gates—a necessity for a universal quantum computation. Here, we adapt results which were obtained by Solenov and Mozyrsky<sup>601</sup> for the case of homogeneous rings with impurities. It results, that a single ring optical lattice with an impurity is described by the following effective Lagrangian [see Eq. (73)]:

$$L = \frac{1}{2U} \dot{\theta}^2 + \frac{J}{N} (\theta - \Phi)^2 - J' \cos \theta. \quad (80)$$

Then, we introduce the canonical momentum  $P$  in a usual way,



**FIG. 45.** Six first energy levels of the reduced system given by the effective potential Eq. (79) as a function of the dimensionless external flux  $\Omega$ . Here  $J' = 0.7J$ ,  $J'' = 0.8J$ ,  $U = 0.5J$ , and  $\theta_1 = -\theta_2$ . Reprinted with permission from Aghamalyan *et al.*, New J. Phys. **18**, 075013 (2016). Copyright 2016, Author(s) licensed under a Creative Commons Attribution 3.0 License.

$$P = \frac{\partial L}{\partial \dot{\theta}} = \frac{1}{U} \dot{\theta}. \quad (81)$$

After performing a Legendre transformation, we get the following Hamiltonian:

$$H = J' \left[ \frac{p^2}{2\mu} - \frac{J}{J'(N-1)} (\theta - \Phi)^2 + \cos \theta \right], \quad (82)$$

where  $\mu = J'/U$  is an effective mass of the collective particle. The quantization is performed by the usual transformation  $P \rightarrow -d/d\theta$ . For  $\delta = J'(N-1)/2J > 1$ , the effective potential in Eq. (82) can be reduced to a double well; for  $\Phi = \pi$ , the two lowest levels of such a double well are symmetric and antisymmetric superpositions of the states in the left and right wells, respectively. The effective Hamiltonian can be written as

$$H \simeq \varepsilon \sigma_z, \quad (83)$$

and the lowest two states are  $|\psi_g\rangle = (0, 1)^T$  and  $|\psi_e\rangle = (1, 0)^T$ . WKB estimate for the energy splitting  $\varepsilon$  of the qubit is given by Eq. (74).

### 1. Single qubit gates

For the realization of single-qubit rotations, we consider the system close to the symmetric double well configuration  $\Phi \simeq \pi$ . In the basis of the two-level system discussed before, the Hamiltonian takes the form

$$H \simeq \varepsilon \sigma_z + \frac{\Phi - \pi}{\delta} \langle \theta \rangle_{01} \sigma_x, \quad (84)$$

where  $\langle \theta \rangle_{01}$  is the off-diagonal element of the phase-slip in the two-level system basis. It is easy to show that spin flip, Hadamard, and phase gates can be realized by this Hamiltonian. For example, a phase gate can be realized by evolving the state through the unitary transformation  $U_z(\beta)$  [tuning the second term of Eq. (84) to zero by adjusting the imprinted flux]

$$U_z(\beta) = \exp(i\varepsilon\tau\sigma_z) = \begin{pmatrix} e^{i\varepsilon\tau} & 0 \\ 0 & e^{-i\varepsilon\tau} \end{pmatrix}. \quad (85)$$

After tuning the gap energy close to zero (adjusting the barrier height of the impurity), we can realize the following rotation

$$U_x(\beta) = \exp(i\alpha\tau\sigma_x) = \begin{pmatrix} \cos \alpha & i \sin \alpha \\ i \sin \alpha & \cos \alpha \end{pmatrix}, \quad (86)$$

where  $\alpha = \Phi - \pi/\delta\langle\theta\rangle_{01}\tau$ . When  $\alpha = \pi/2$  and  $\alpha = \pi/4$ , the NOT and Hadamard gates are, respectively, realized.

### 2. Two-qubit coupling and gates

The effective dynamics for two coupled qubits, each realized as single ring with localized impurity (as in Fig. 50), is governed by the Lagrangian

$$L = \sum_{a,b} \frac{1}{2U} \dot{\theta}_a^2 + \left[ \frac{J}{2(N-1)} (\theta_a - \Phi_a)^2 - J' \cos(\theta_a) \right] - \tilde{J}'' \cos \left[ \theta_a - \theta_b - \frac{N-2}{N} (\Phi_a - \Phi_b) \right], \quad (87)$$

where  $J''$  is the Josephson tunneling energy between the two rings. When  $\Phi_a = \Phi_b = \Phi$  and  $J'' \ll J'$ , the last term reduces to  $-J''(\theta_a - \theta_b)^2/2$  and the Lagrangian takes the form

$$L = J' \left[ \sum_{a,b} \frac{1}{2J'U} \dot{\theta}_a^2 + \left[ \frac{J}{2J'(N-1)} (\theta_a - \Phi_a)^2 - \cos(\theta_a) \right] + \frac{J''}{J'} \frac{(\theta_a - \theta_b)^2}{2} \right]. \quad (88)$$

By applying the same procedure as in Sec. X B 1, we obtain the following Hamiltonian in the eigenbasis of the two-level systems of rings  $a$  and  $b$ ,

$$H = H_a + H_b + \frac{J''}{J'} \sigma_x^1 \sigma_x^2 \langle \theta \rangle_{01}^2, \quad (89)$$

$$H_x = \varepsilon \sigma_z^x + \left( \frac{\Phi - \pi}{\delta} + \frac{J''\pi}{J'} \right) \langle \theta \rangle_{01} \sigma_x^x. \quad (90)$$

From this equations, it follows that qubit-qubit interactions can be realized using our setup. If we choose the tuning  $\varepsilon \rightarrow 0$  and  $\Phi \rightarrow \pi - \delta J''\pi/J'$ , the natural representation of a  $(\text{SWAP})^x$  gate<sup>602</sup> can be obtained,

$$U(\tau) = \exp \left[ -i \frac{J''}{J'} \sigma_x^1 \sigma_x^2 \tau \right], \quad (91)$$

where  $\alpha = \tau J''/J'$ . A CNOT gate can be realized by using two  $\sqrt{\text{SWAP}}$  gates.<sup>602</sup> It is well known that one qubit rotations and a CNOT gate are sufficient to implement a set of universal quantum gates.<sup>603</sup>

### C. Readout of atomtronic qubits

For a controlled quantum system, it is essential to be able to read out the state of a prepared quantum state. For the atomtronic qubit, one can determine the properties of the qubit by reading out the current of the atoms. The existence of the atomic current flowing in AQUID can be detected by standard time-of-flight measurement of the ring condensate.<sup>14</sup> A more in-depth analysis can be performed by interfering the ring condensate with a second condensate confined in the center of a ring. This condensate sets a phase reference for the phase winding of the ring condensate. By *in-situ* measurement of the two interfering condensates, the self-heterodyne detection of the phase of the wave function is realized. For weakly interacting continuous ring systems, where no entanglement is present, both the orientation and the intensity of the current states have been detected.<sup>149,392,604,605</sup>

For atomtronic qubits, this detection scheme has to be applied to the case of ring lattices with stronger interactions. This has been studied in Ref. 590, and the key results are reviewed below.

#### 1. Interferometric detection of the current states

To read out the direction and the intensity of the current in the ring lattice, an approach originally carried out by the Maryland and



Paris groups to map-out the circulating states in continuous ring-shaped condensates can be applied.<sup>16,142,161,392,604</sup> Accordingly, the ring condensate is made to interfere with another condensate at rest, located at the center, fixing the reference for the phase of the wavefunction. The combined wavefunction evolves in time, interferes with itself and finally is measured. The number of spirals gives the total number of rotation quanta.

In the actual experiment, the condensate is imaged through *in situ* measurements. In this way, the current direction and magnitude is well visible as a spiral pattern. The position of the spirals depends on the relative phase between ring and the central condensate. In a single experimental run, the spirals will be visible for a condensate with the high number of particles. However, if the number of particles is low or the atom imaging is inefficient, one has to average over multiple shots and take expectation values, which experimentally corresponds to take averaged results over many experimental runs. However, every realization of the experiment has a random phase in the phase of the spirals, which is averaged out over many repetitions. As the relative phase between ring and central condensate is determined randomly upon measurement, the expectation value of the density operator will average over different realizations of the spiral interference pattern, washing out the information on the current configuration structure. However, as we show below, the information about the spirals can be recovered using density-density correlations.

The expansion dynamics is modeled with the Bose-Hubbard model. The ring wavefunction is calculated by solving the ground state of the Bose-Hubbard Hamiltonian, while the central condensate is simply a single decoupled site with  $N_c$  particles. The dynamics of the density  $\hat{n}(\mathbf{r}, t) = \hat{\psi}^\dagger(\mathbf{r}, t)\hat{\psi}(\mathbf{r}, t)$  is initialized assuming that the bosonic field operator of the system is  $\hat{\psi}(\mathbf{r}) = \sum_n w_n(\mathbf{r})\hat{a}_n$ , where  $w_n(\mathbf{r})$  are a set of Wannier functions forming a complete basis.<sup>606,607</sup> In our calculation, we approximate the full basis for wave functions living in the ambient space on which the condensate expands with the set of Wannier functions composed of Gaussians peaked at the ring lattice sites and at its center (the Gaussian approximation for the Wannier functions is a well verified approximation for single site wavefunctions—see Refs. 608 and 609) For the free evolution (we are indeed in a dilute limit), we assume that each particle at site  $n$  expands in two dimensions as

$$w_n(\mathbf{r}, t) = \frac{1}{\sqrt{\pi}} \frac{\sigma_n}{\sigma_n^2 + \frac{i\hbar t}{m}} e^{-((\mathbf{r}-\mathbf{r}_n)^2/2(\sigma_n^2 + i\hbar t/m))}, \quad (92)$$

where  $\sigma_n$  is the width of the condensate located at the  $n$ -th site. The dynamics of the condensates is then approximated as  $\hat{\psi}(\mathbf{r}, t) = \sum_n w_n(\mathbf{r}, t)\hat{a}_n$ . We observe that such an approximation works well in the situations in which the optical lattice is assumed to be sufficiently dense in the space in which the condensate is released (as in the release from large three dimensional optical lattices).

To observe the interference, and thus the qubit properties, using averaging over multiple shots, the interference pattern is measured with higher order density-density correlations. We calculate the density-density covariance<sup>610–613</sup>

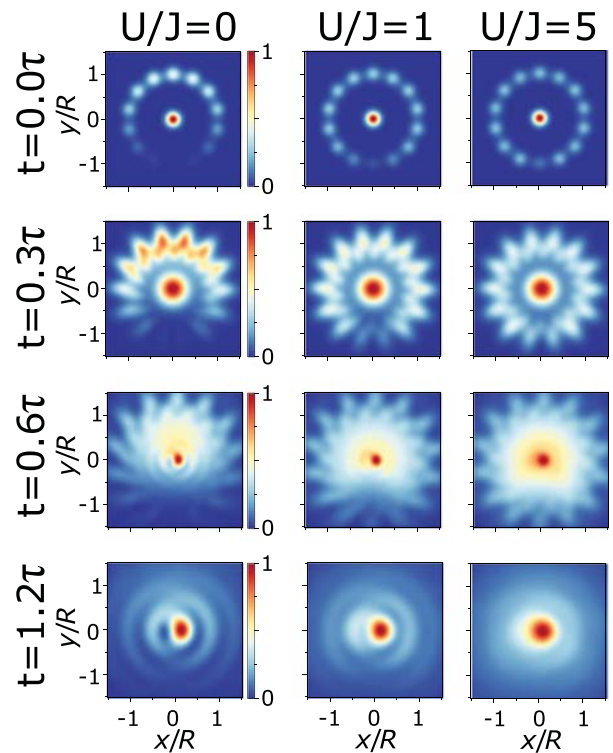
$$\text{cov}(\mathbf{r}, \mathbf{r}', t) = \langle \hat{n}(\mathbf{r}, t)\hat{n}(\mathbf{r}', t) \rangle - \langle \hat{n}(\mathbf{r}, t) \rangle \langle \hat{n}(\mathbf{r}', t) \rangle. \quad (93)$$

We also define the root of the density covariance, which has the same unit as the density to improve the contrast of the measured interference pattern,

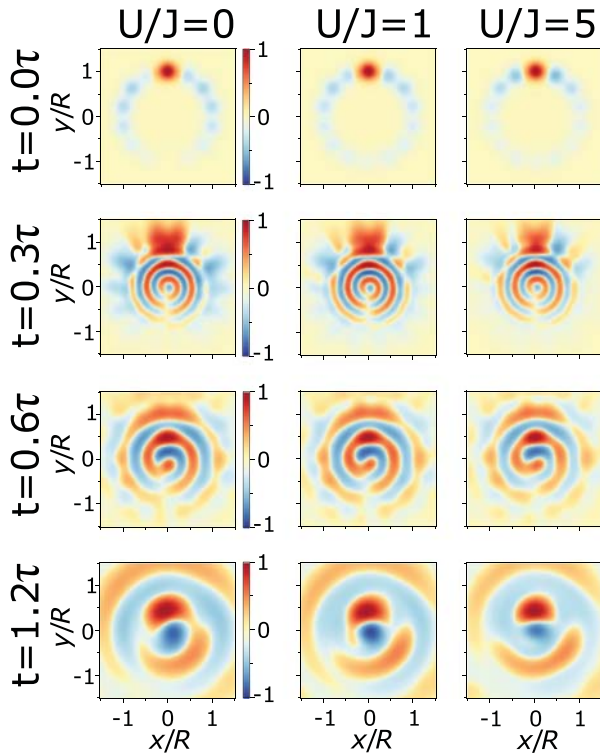
$$\sigma(\mathbf{r}, \mathbf{r}', t) = \text{sgn}(\text{cov}(\mathbf{r}, \mathbf{r}', t)) \sqrt{|\text{cov}(\mathbf{r}, \mathbf{r}', t)|}. \quad (94)$$

First, we plot the expectation value of the density of expanded atoms for different values of interaction at the degeneracy point  $\Omega = 1/2$  in Fig. 46. The density of expanded atoms at longer times has some characteristic features depending on the interaction. For interaction energy smaller than the potential barrier, the center shows a characteristic bright and dark spot. For stronger interaction, it becomes a single, blurred spot. At the degeneracy point, we observe a superposition of counter-flowing current states. Interaction modifies the many-body entanglement, which changes the characteristic time-of-flight pattern. After a long enough free expansion, the atom density assumes the initial momentum distribution. However, it is difficult to read out the exact state of the current as the characteristic spirals are not visible in the expectation values of the density.

Next, we show the density-density covariance  $\sigma(\mathbf{r}, \mathbf{r}')$  in Fig. 47. A clear spiral pattern emerges here. In this case, a step in the spirals at the weak link site (here at the center bottom) is clearly visible for intermediate times. This indicates the appearance of a superposition of two winding numbers. Although the interferometric pictures can look



**Fig. 46.** Density of expanding atoms at times  $t = 0, 0.3\tau, 0.6\tau, 1.2\tau$ , with  $\tau = mR\sigma_r/\hbar$ . From left to right:  $U=0$ ,  $U/J=1$ ,  $U/J=5$ . Flux  $\Omega = \frac{1}{2}$  at the degeneracy point. At intermediate time, we observe some spiral-like structure at the edges. This is not the interference with the central condensate, but a residue of the ring lattice interfering with itself. Calculated using Bose-Hubbard model, no interaction during expansion. Data in color and normalized to one. Ring has 7 particles,  $M = 14$  ring sites, ring radius  $R$ . Width of central and ring cloud is  $\sigma_r = 2R/L$  and potential barrier  $\Lambda = J$ , 25% of atoms in central condensate. Barrier at  $x=0$ ,  $y=-R$ . Reprinted with permission from Haug *et al.*, Phys. Rev. A **97**, 013633 (2018). Copyright 2018, American Physical Society.



**FIG. 47.** Root of density-density covariance  $\sigma(\mathbf{r}, \mathbf{r}') = \{0, R/2\}$  of expanding atoms with flux  $\Omega = \frac{1}{2}$  at the degeneracy point. The discontinuity in the bottom of the spirals at intermediate times  $t = 0.3\tau$  and  $t = 0.6\tau$  shows that the ring condensate is in a superposition of zero and one rotation quantum. Same parameters as Fig. 46. Reprinted with permission from Haug *et al.*, Phys. Rev. A **97**, 013633 (2018). Copyright 2018, American Physical Society.

similar, different interactions lead to current states that may be very different in nature. For  $U = 0$ , the current is in a nonentangled superposition state, whereas for interaction  $U = J$  in a highly entangled NOON state.

Below, we shall see how additional information on the states can be grasped analyzing the noise in the momentum distribution of the ring condensate. Indeed, the noise for zero momentum strongly depends on the specific entanglement between the clockwise and anti-clockwise flows. In the case of an entangled cat state, all atoms have together either zero or one momentum quanta. A projective measurement will collapse the wavefunction to either all atoms in the zero or one momentum state. Averaging over many repeated measurements will result in erratic statistics of the measurements. In contrast, in nonentangled single-particle superpositions, each particle has independently either zero or one momenta quanta. A single projective measurement will result in on average half the atoms having zero and half the atoms having one rotation quantum. Therefore, fluctuations averaged over many measurements will be low. We define the noise of the momentum distribution,

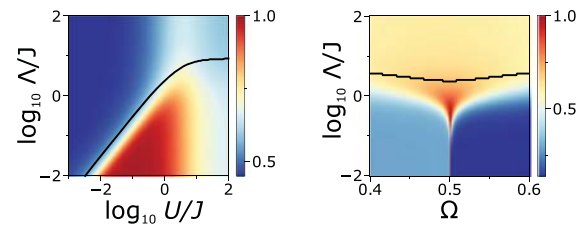
$$\sigma_k(\mathbf{k}) = \sqrt{\langle \hat{n}(\mathbf{k}) \hat{n}(\mathbf{k}) \rangle - \langle \hat{n}(\mathbf{k}) \rangle \langle \hat{n}(\mathbf{k}) \rangle}. \quad (95)$$

Having in mind a time-of-flight experiment, the optimal point to measure the noise is at  $\mathbf{k} = 0$ , as at this point the density is maximal for

zero rotation quanta, and zero for one or more rotation quanta. We plot the noise of the time-of-flight image at  $\mathbf{k} = 0$  without a central condensate in Fig. 48. First, the interaction  $U$  and weak link  $\Lambda$  is plotted in Fig. 48(a). We see that the momentum noise is minimal in the parameter regime  $U/J \ll 1$  and  $\Lambda/J > cU/J$ , where  $c$  is some constant, which corresponds to the mean-field limit. As soon the interaction becomes larger than the energy gap induced by the potential barrier, the noise increases. Here, entangled phase winding states of zero and one winding quantum appear. For the large interaction, the noise decreases again, however, remains higher than in the mean-field regime. With the increasing interaction, we can define three regimes of entanglement.<sup>614</sup> At the degeneracy point  $\Omega = \frac{1}{2}$ , for interaction smaller than the energy gap created by the weak link, we observe one-particle superposition states  $|\Psi\rangle \propto (|l=0\rangle + |l=1\rangle)^N$ , where  $N$  is the number of particles and  $l$  is the angular momentum of the atom. This regime is well described by the Gross-Pitaevskii equation. Here, the noise at  $\mathbf{k} = 0$  is minimal and is given by  $\sigma_k^{\text{GP}}(\mathbf{k} = 0) \propto \sqrt{N}/4$ . When the interaction and the weak-link energy gap is on the same order, the near-degenerate many body states mix and entangled NOON states are formed  $|\Psi\rangle \propto |l=0\rangle^N + |l=1\rangle^N$ . The noise is maximal and given by  $\sigma_k^{\text{NOON}}(\mathbf{k} = 0) \propto N/4$ . The ratio of the minimal and maximal noise is  $\sqrt{N}$ . Thus, with the increasing particle number, the superposition and entangled states are clearer to distinguish. The increasing interaction further will fermionize the system. With the interaction, angular momentum of each atom individually is not conserved, however, the center of mass angular momentum  $K$  of the whole condensate is. Then, the ground state is a superposition of  $|\Psi\rangle \propto |K=0\rangle + |K=N\rangle$ .

Next, the momentum noise is plotted against applied flux  $\Omega$  in Fig. 48(b). Due to the two level system effective physics, the noise in the time of flight of the ring condensate is particularly pronounced at the degeneracy points. This phenomenon allows one to detect the degeneracy point in the ring condensate, without resorting the heterodyne detection protocol. The noise is maximal at the degeneracy point, when the barrier and interaction are on the same order. Changing the flux away from the degeneracy point decreases the noise.

Further information can be identified by looking at the density at the site of the weak link. For the zero on-site interaction, the site at the potential barrier is always depleted at the degeneracy point for any value of potential barrier strength. However, when the interaction



**FIG. 48.** Momentum noise  $\sigma_k(\mathbf{k} = 0)$  (in color, normalized to one) plotted for potential barrier  $\Lambda$  against a) on-site interaction  $U$  ( $\Omega = \frac{1}{2}$ ) and b) flux  $\Omega$  ( $U/J = 1$ ). Momentum noise is extracted from time-of-flight image after long expansion. Only ring is expanded, without central condensate. Black line shows the critical point where depletion at the potential barrier is 1% of the average particle number per site. Above the line the potential barrier site is depleted. Other parameters are  $M = 11$  ring sites and 5 particles. Reprinted with permission from Haug *et al.*, Phys. Rev. A **97**, 013633 (2018). Copyright 2018, American Physical Society.

exceeds a critical value, particles start occupying the site.<sup>14</sup> This is plotted as a black line in Fig. 48. For small interaction, the critical value has a linear relationship between  $U$  and  $\Lambda$ .<sup>14</sup> The filling of the potential barrier site indicates the onset of entanglement between different flux quanta. The depletion factor can be measured by a lattice-site resolved absorption measurements.

#### D. Experiment realization of the ring-lattice potential with weak links

In this section, we provide the experimental details for the realization of ring-lattice potentials with weak-links. Among the different architectures, the focus is on the structure that can be relevant for the construction of two level quantum systems.

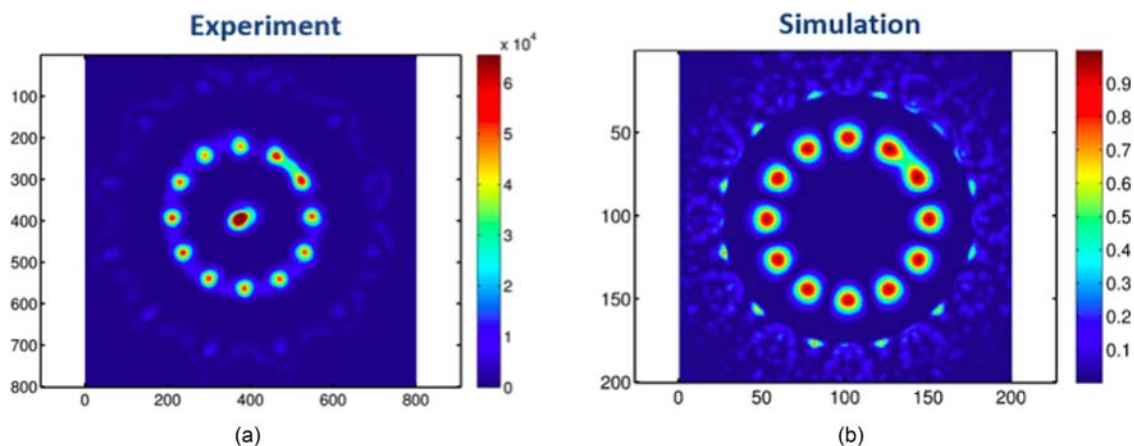
##### 1. A ring lattice with single weak link

The optical potential was created with a liquid crystal on a silicon spatial light modulator (PLUTO phase only SLM, Holoeye Photonics AG), which imprints a controlled phase onto a collimated laser beam from a 532 nm wavelength diode pumped solid state (DPSS) laser. The SLM acts as a programmable phase array and locally modifies the phase of an incoming beam. Diffracted light from the computer generated phase hologram then forms the desired intensity pattern in the focal plane of an optical system (doublet lens,  $f = 150$  mm). The resulting intensity distribution is related to the phase distribution of the beam exiting the SLM by Fourier transform. The calculation of the required SLM phase pattern (kinoform) has been carried out using an improved version of the mixed-region-amplitude-freedom (MRAF) algorithm<sup>59,615</sup> with angular spectrum propagator. This allows one to numerically simulate the wavefront propagation in the optical system without resorting to paraxial approximation. A region outside the desired ring lattice pattern (noise region) is dedicated to collect unwanted light contributions resulting from the MRAF algorithm's iterative optimization process. This can be seen in the measured intensity pattern in Fig. 49 as concentric, periodic structures surrounding the ring-lattice and can be filtered out by an aperture.

The ring-lattice potential shown in Figs. 49 and 50 can be readily scaled down from a radius of  $\sim 90$  to  $5 - 10 \mu\text{m}$  by using a  $50\times$  microscope objective with  $\text{NA} = 0.42$  numerical aperture (Mitutoyo  $50\times$  NIR M-Plan APO) as the focusing optics for the SLM beam and with  $\lambda_2 = 830$  nm light, suitable for trapping rubidium atoms. Accounting for the limited reflectivity and diffraction efficiency of the SLM, scattering into the noise region and losses in the optical system only about 5% of the laser light contributes to the optical trapping potential. However, this is not a limiting factor for small ring-lattice sizes in the tenth of micrometer range as discussed here where  $\sim 50$  mW laser power is sufficient to produce well depths of several  $E_{\text{rec}}$ . The generated structures are sufficiently smooth with a measured intensity variation of 4.5% rms to sustain persistent flow-states.<sup>67</sup> The barrier height can be dynamically modified at a rate up to 50 ms per step with an upper limit imposed by the frame update rate of the SLM LCD panel (60 Hz).

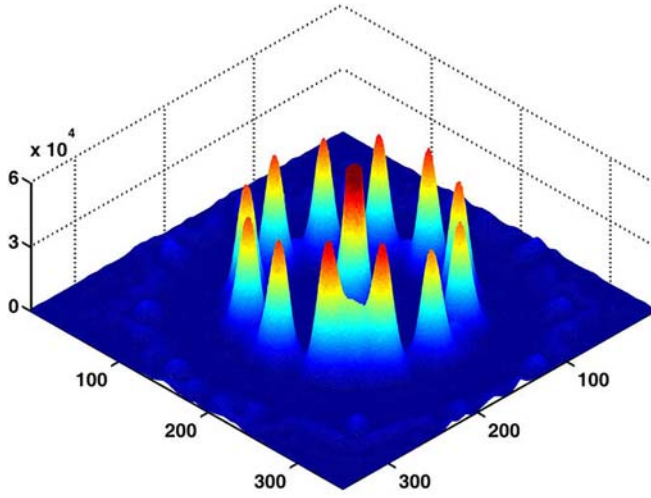
##### 2. Experiment realization of the ring-lattice potential with three weak links

We produce the optical potential using a spatial light modulator (Holoeye Photonics AG, PLUTO-NIR II), SLM. A collimated Gaussian beam, of 8 mm diameter, is reflected from the SLM's surface forming an image through a  $f = 200$  mm lens. The light is then split into the two sides of our system with 10% of the light in the "monitoring" arm and 90% into the "trapping" arm used to create a red-detuned dipole trapping potential for a gas of  $\text{Rb}^{87}$  atoms. A Ti:sapphire laser (Coherent MBR-110) produces a 1 W, 828 nm beam, which is spatially filtered and collimated, before reflection on the SLM. To produce the trapping potential, the SLM's kinoform is imaged through a  $4f$  lens system reducing the beam size to 3 mm diameter and focused through a  $50\times$  microscope objective with a 4 mm focal distance and a numerical aperture of  $\text{NA} = 0.42$  (Mitutoyo  $50\times$  NIR M-Plan APO). The monitoring arm of the system creates an image of the potential through a  $10\times$  infinity-corrected microscope objective focused on a CCD camera (PointGrey FL3-GE-13S2M-C). The CCD camera views, therefore, an enlarged image of the optical potential.



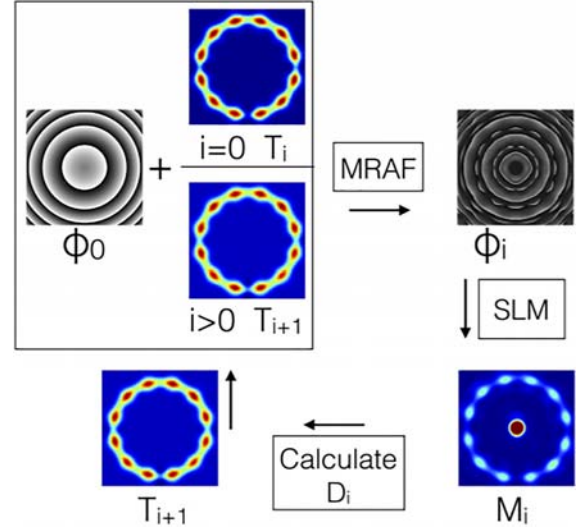
**FIG. 49.** Simulation (a) and experimentally realized (b) intensity distribution of a ring-lattice with a weak link between two lattice sites. Reprinted with permission from Aghamalyan, Ph.D. dissertation, Singapore, 2015.





**FIG. 50.** Experimental realization of a ring-lattice potential with an adjustable weak link. Measured intensity distribution with an azimuthal lattice spacing of  $28 \mu\text{m}$  and a ring radius of  $88 \mu\text{m}$ . The central peak is the residual zero-order diffraction. The size of the structure is scalable and a lower limit is imposed by the diffraction limit of the focusing optics. Reprinted with permission from Amico *et al.*, Sci. Rep. **4**, 4298 (2014). Copyright 2014, Author(s) licensed under a Creative Commons Attribution 3.0 License.

To increase the accuracy of the output potential, we use the computationally generated kinoform and produce an image of the optical potential in the monitoring arm of our system and use this as a further source of feedback to the MRAF algorithm. Our method is broadly similar to Bruce *et al.*,<sup>616</sup> however it is specialized for producing ring-lattices. Figure 51 shows a flow chart of our improved algorithm. In the first step, the target image,  $T_i$ , and the initial phase,  $\phi_0$ , are loaded as an input to the MRAF code. This runs for 20 iterations (this was found to be sufficient to get good convergence in most cases) and outputs a phase kinoform,  $\phi_i$ . The kinoform is now applied to the SLM and an image recorded on the camera in the monitoring arm of our system,  $M_i$ . The discrepancy,  $D_i$ , between the original target and the measurement is calculated and used to form an updated target  $T_{i+1}$ . Here, our algorithm differs from Ref. 616 as we take the discrepancy to be  $D_i = -(M_i^2 + T_0^2)/2T_0$ . Also, we do not take into account the whole image, and the discrepancy is calculated by comparing the maxima and minima around the azimuthal, 1D, profile of the lattice to the target profile. The targets maxima and minima are then adjusted with  $T_{i+1} = T_i + \alpha D_i$ , where  $\alpha$  is a problem specific feedback gain and  $i$  is the iteration number. The process now repeats with  $\phi_0$  and  $T_{i+1}$  as the inputs to the MRAF code. The feedback gain,  $\alpha$ , is set to be 0.3 to ensure a quick convergence, and this process iterates 30 times. At this point, the algorithm is complete and the best image from the set  $M$  is selected that minimizes the discrepancy below 2%. With this method, we produce the ring-lattice potential shown in Fig. 52 (left) that on the trapping side of our apparatus creates a scaled-down lattice with a radius of  $5\text{--}10 \mu\text{m}$  with more than sufficient power to trap ultracold atoms. On the right of Fig. 52, the azimuthal profile around the ring lattice is shown. The red curve indicates the profile on the first iteration of the feedback loop. After five iterations (blue curve), the algorithm has converged significantly toward the original target (solid line).



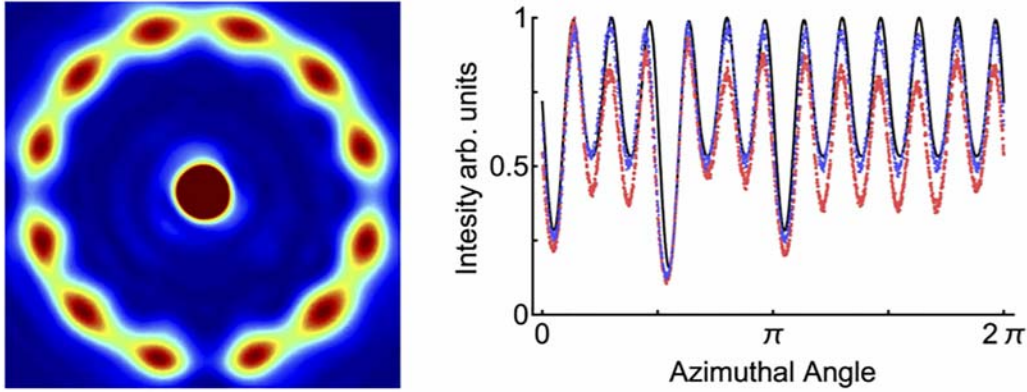
**FIG. 51.** Our feedback algorithm. Starting at the top left the initial phase and target are used in the MRAF code. This generates the phase guess,  $\phi_i$ , which is uploaded to the SLM and an image captured by the CCD camera,  $M_i$ . This is used to calculate the discrepancy between the image and the original target, and a new target  $T_{i+1}$  is created. The loop then repeats. Reprinted with permission from Aghamalyan *et al.*, New J. Phys. **18**, 075013 (2016). Copyright 2016, Author(s) licensed under a Creative Commons Attribution 3.0 License.

## E. Setup for adjustable ring-ring coupling

In this section, proposal for experimental realization of ring lattices with tunable distance between the rings is suggested by utilizing the SLM technique.<sup>59,615</sup> With a SLM arbitrary, optical potentials can be produced in a controlled way only in a  $2d$ -plane—the focal plane of the Fourier transform lens—making it challenging to extend and up-scale this scheme to 3d trap arrangements. The experiment, however, showed (see Fig. 53) that axially the ring structure potential remains almost undisturbed by a translation along the beam propagation axis of  $\Delta z = \pm 2.2 \cdot R$ , where  $R$  denotes the ring-lattice radius. The ring-lattice radius is only weakly affected by an axial shift along  $z$  and scales with  $\Delta R/R = 0.0097 \cdot z$ , where  $z$  is normalized to the ring-lattice radius. For larger axial shifts from the focal plane, the quality of the optical potential diminishes gradually. Based on our measurements, this would allow implementation of ring-lattice stacks with more than ten rings in a vertical arrangement, assuming a stack separation comparable to the spacing between two adjacent lattice sites. Propagation invariant beams may allow a potentially large number of rings to be vertically arranged.<sup>532</sup>

In addition to making the inter-ring dynamics strictly one dimensional, the lattice confinement provides the route to the inter-rings coupling.

To allow controlled tunneling between neighboring lattice along the stack, the distance between the ring potentials needs to be adjustable in the optical wavelength regime (the schematics in Fig. 54 can be employed). A trade-off between high tunneling rates (a necessity for fast gate operations) and an efficient readout and addressability of individual stack sites needs to be analyzed. Increasing the lattice stack separation after the tunneling interaction has occurred well above the



**FIG. 52.** Left: Final image of the ring lattice after completion of the feedback algorithm. Right: Azimuthal Profile. The solid line plots the target profile. This is compared to the result after the 1st and 5th iteration of the feedback algorithm (red and blue lines, respectively). Reprinted with permission from Aghamalyan *et al.*, New J. Phys. **18**, 075013 (2016). Copyright 2016, Author(s) licensed under a Creative Commons Attribution 3.0 License.

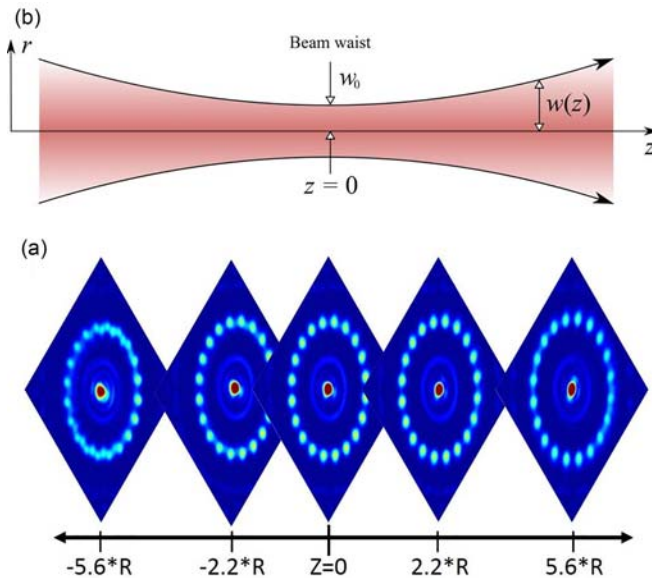
diffraction limit while keeping the atoms confined, optical detection, and addressing of individual rings becomes possible.

This arrangement produces equal, adjustable ring-ring spacing between individual vertical lattice sites and can, therefore, not readily be used to couple two two-ring qubits to perform two-qubit quantum-gates. The SLM method, however, can be extended to produce two ring-lattices in the same horizontal plane separated by a distance larger

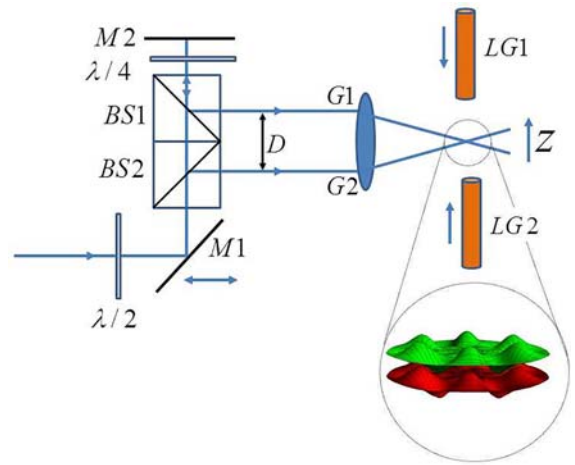
than the ring diameter. The separation between these two adjacent rings can then be programmatically adjusted by updating the kinoform to allow tunneling by mode overlap.<sup>618</sup> Combined with the adjustable vertical lattice (shown in Fig. 54), this would allow, in principle, two-ring qubit stacks to be circumferential tunnel-coupled to form two-qubit gates.

#### F. First experimental demonstration of the interference of atomtronic currents

A key ingredient to realize the atomtronic qubit is the interference of currents that make up the qubit dynamics. Controlling and interfering currents *in situ* of a cold atom circuit are a challenging task,



**FIG. 53.** Effect of an axial translation on the ring lattice potential. (a) Ring lattice intensity distribution measured at various positions along the beam propagation axis around the focal plane ( $Z=0$ ). Note that the initial beam, phase modified by the SLM, is not Gaussian any more. The optical potential remains undisturbed by a translation of 2.2 times the ring-lattice radius centered around the focal plane ( $Z=0$ ). Here  $R$  designates the ring-lattice radius of  $87.5 \mu\text{m}$ . (b) This is in contrast to a Gaussian laser beam which exhibits a marked dependence on the axial shift from the focal plane where the beam waist  $w(z)$  scales with  $\sqrt{1 + (z/z_0)^2}$  and Rayleigh range  $z_0$ . Reprinted with permission from Amico *et al.*, Sci. Rep. **4**, 4298 (2014). Copyright 2014, Author(s) licensed under a Creative Commons Attribution 4.0 License.



**FIG. 54.** Proposed setup for the ring-ring coupling. Two Gaussian laser beams of wavelength  $\lambda$  and distance  $D$ , pass through a lens and interfere in the focal plane ( $f$  is the focal length). The distance  $D$  can be easily controlled by moving the mirrors. The distance between the fringes is a function of  $1/D$ .<sup>617</sup> The resulting Gaussian laser beam with wave vector  $k_G = 2\pi D/(\lambda f)$ , then, interferes with two counter propagating Laguerre-Gauss laser beams of amplitude  $E_0$ . The inset shows the ring lattice potentials separated by  $d = \lambda f/D$ . Here,  $l=6$  and  $p=0$ . Reprinted with permission from Aghamalyan *et al.*, Phys. Rev. A **88**, 063627 (2013). Copyright 2013, American Physical Society.

as it requires high coherence and control in the system. Recently, interference of currents in an AQUID has been realized experimentally for the first time, which is reviewed in this section.<sup>15</sup> The interference can be revealed by a periodic modulation of the critical current with applied flux in an atomtronic ring interrupted by two weak links. Note that the interference has been achieved for a dilute Bose–Einstein condensate, which can be described within the mean-field limit as a classical wave equation. While entanglement that is critical for atomic qubits cannot be demonstrated within this experiment, it is nonetheless a first step toward establishing the ingredients for atomic qubits based on superposition of currents.

The periodic modulation of the critical current can be understood by calculating the total current within a model of the atomtronic SQUID based on quantum phase-controlled Josephson junction currents and a toroidal trap geometry [Fig. 55(a)]. The total current is the result of interference of the two Josephson junction currents, given by

$$I_1 = \frac{1}{2}(I_t + I_j) = I_c \sin(\phi_1), \quad (96)$$

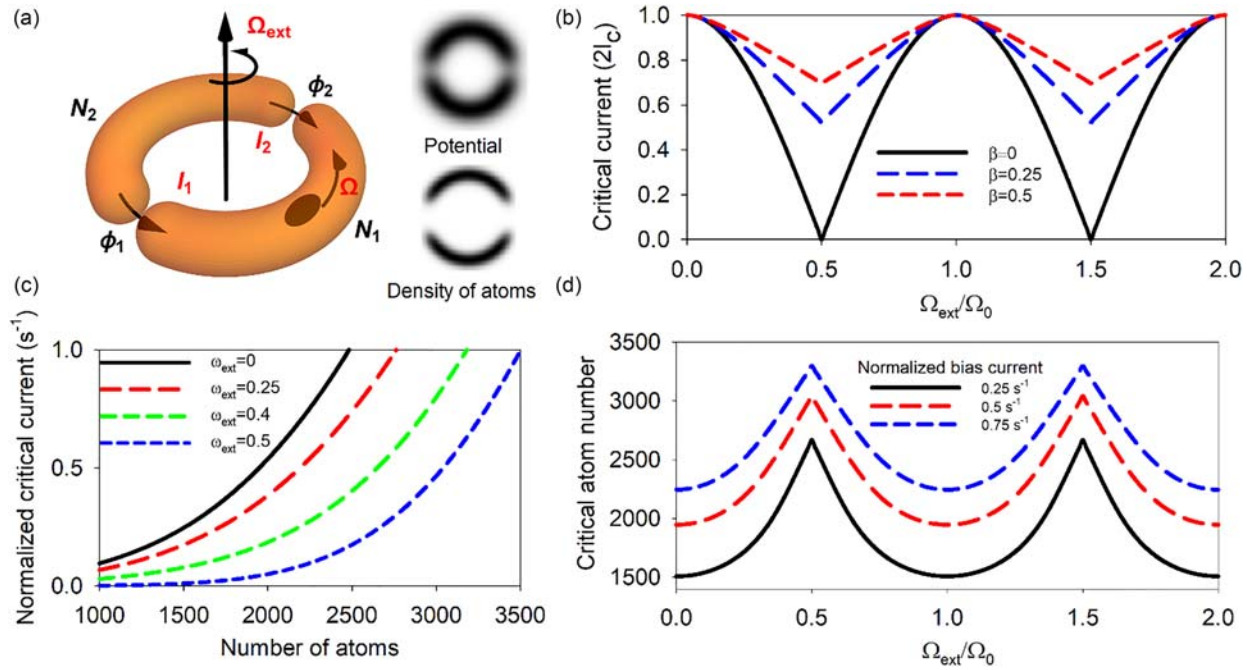
$$I_2 = \frac{1}{2}(I_t - I_j) = I_c \sin(\phi_2), \quad (97)$$

where  $I_c$  is the critical current of atoms,  $I_t$  is the total current, and  $I_j$  is the circulating current around the atomtronic SQUID. Because of the

toroidal geometry and single valuedness of the wavefunction describing the atoms, the phases should satisfy  $\phi_1 - \phi_2 + 2\pi\omega = 2\pi n$ , where  $\omega = \Omega/\Omega_0$ , with  $\Omega$  being the rotation rate of atoms and  $n$  is an integer. The rotation rate of the atoms can be shown to be

$$\omega = \omega_{\text{ext}} + \beta_{\text{atom}} \frac{I_j}{I_c}, \quad (98)$$

where  $\omega_{\text{ext}} = \Omega_{\text{ext}}/\Omega_0$ ,  $\Omega_{\text{ext}}$  is the external rotation rate of the atomtronic SQUID,  $\beta_{\text{atom}} = 2\pi I_c/N\Omega_0$ , and  $N$  is the total number of atoms. This equation for the rotation rate of atoms can be derived from the relation between the circulating current and the movement of atoms relative to the Josephson junctions. The parameter  $\beta_{\text{atom}}$  is analogous to the screening parameter in the conventional SQUID and can be thought as proportional to the “inductance” that induces the deviation of the rotation rate of atoms from the imposed external rotation rate of the atomtronic SQUID. Equations (1)–(4) are equivalent to those of a DC SQUID,<sup>619</sup> reflecting the fact that the fundamental underlying physics of a double junction atomtronic SQUID and a DC SQUID is the same. In the limit of  $\beta_{\text{atom}} = 0$  (for example, when  $I_c \approx 0$  with much higher barrier height), we can analytically calculate the total currents  $I_t = 2I_c \cos(\pi\omega_{\text{ext}}) \sin(\phi_1 - \pi\omega_{\text{ext}})$ . Thus, the critical current is  $|2I_c \cos(\pi\omega_{\text{ext}})|$ , which establishes a clear modulation of the critical currents by tuning  $\omega_{\text{ext}}$  with a period of  $\Omega_0$ . With finite  $\beta_{\text{atom}}$ ,



**FIG. 55.** Calculation of the periodic modulation of the critical current. (a) Schematic of a double junction atomtronic SQUID. The atomtronic SQUID was created by scanning a single 834 nm laser beam with 1.7  $\mu\text{m}$  waist and the barrier full width at half maximum (FWHM) was 2.1  $\mu\text{m}$ .  $\Omega_{\text{ext}}$  is the rotation rate of the atomtronic SQUID and  $\Omega$  is the rotation rate of atoms.  $\phi_1$  and  $\phi_2$  are the phase differences across the Josephson junctions,  $I_1$  and  $I_2$  are Josephson junction currents, and  $N_1$  and  $N_2$  are numbers of atoms in each half. Arrows represent the movement of the junctions. The calculated potential of the atomtronic SQUID and the density of atoms are shown for the radius of 3.85  $\mu\text{m}$ . (b) Critical current as a function of  $\Omega_{\text{ext}}/\Omega_0$  calculated for different values of  $\beta_{\text{atom}}$ . (c) Normalized critical currents ( $2I_c/N$ ) where  $I_c$  is the critical current and  $N$  is the total number of atoms as a function of the number of atoms with different  $\omega_{\text{ext}}$  for the atomtronic SQUID with 3.85  $\mu\text{m}$  radius.  $\beta_{\text{atom}}$  varies with the number of atoms and the critical current. For each number of atoms,  $\beta_{\text{atom}}$  was calculated to find the variation of the normalized critical current. (d) Modulation of the critical atom number as a function of  $\Omega_{\text{ext}}/\Omega_0$  for three different normalized bias currents with the 3.85  $\mu\text{m}$  radius atomtronic SQUID. Reprinted with permission from Ryu *et al.*, Nat. Commun. 11, 3338 (2020). Copyright 2020, Author(s) licensed under a Creative Commons Attribution 4.0 License.



we can numerically calculate the critical current, and the periodic modulation amplitude decreases with the increasing  $\beta_{\text{atom}}$ , as can be seen in Fig. 55(b). By using the calculated modulation in Fig. 55(b), the expected periodic modulation of the critical current in an atomtronic SQUID can be calculated with the Gross–Pitaevskii equation (GPE) in 2D. Figure 55(c) shows the normalized critical current, which is the critical current of atoms normalized to the number of atoms  $2I_c/N$ , as a function of the number of atoms for the different rotation rates of the atomtronic SQUID. For a fixed number of atoms, the normalized critical current shown in Fig. 55(c) modulates with the rotation rate. However, it is very difficult to experimentally observe this modulation because of the strong dependence of the normalized critical current on the number of atoms and the difficulty in producing a BEC with the same number of atoms consistently. Instead of a fixed number of atoms, a fixed normalized bias current was used, generated by moving Josephson junctions with a fixed speed. When the rotation rate changes, the critical atom number—which is the number of atoms at the transition from DC to AC Josephson effect with the chosen normalized bias current—modulates periodically, as shown in the GPE calculation of Fig. 55(d).

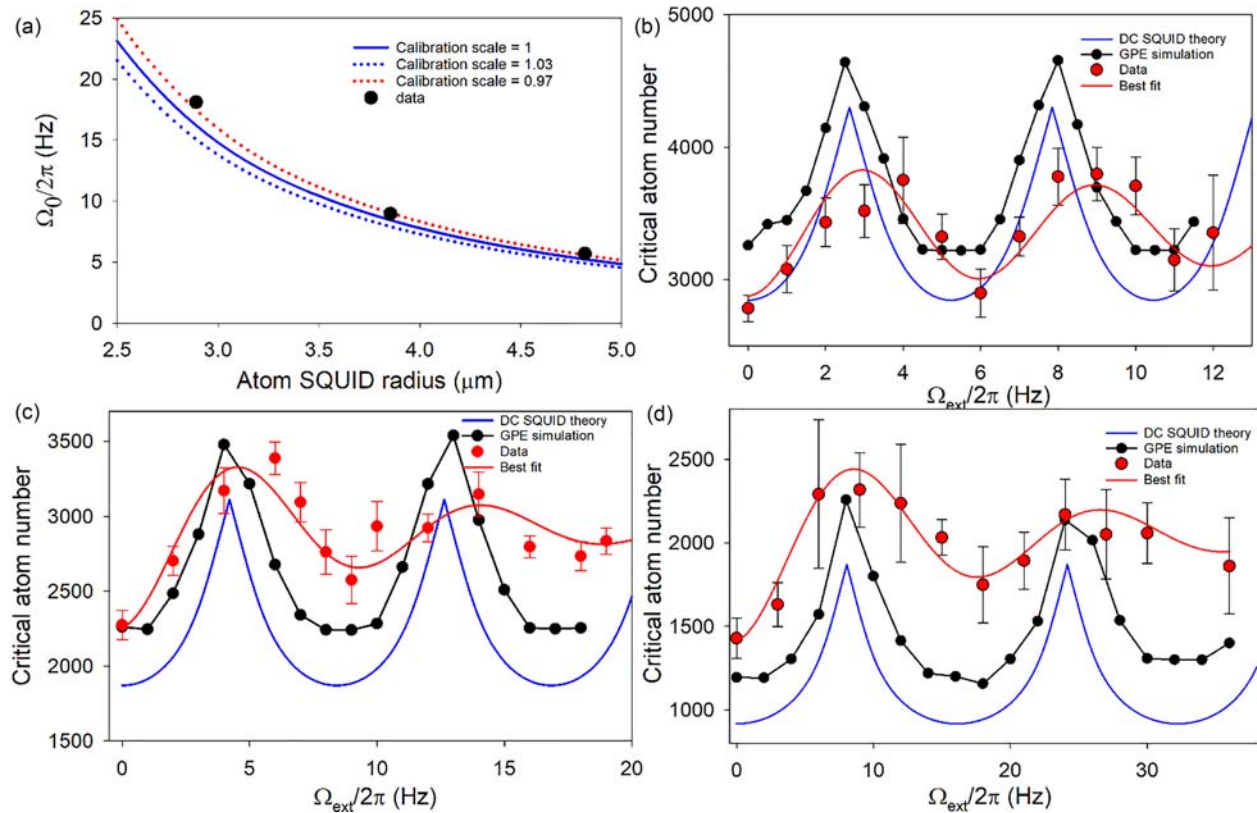
The theoretical prediction of the modulation of the critical current (measured using the critical atom number) is plotted in Fig. 56. The experimental values clearly show the characteristic modulations

of the critical current with flux, revealing the interference of currents in the AQUID.

### G. Concluding remarks and outlook

In this section, we have introduced atomtronic qubits constructed with neutral atomic currents flowing in ring-shaped optical lattice potentials interrupted by few weak links, which give rise to the Atomtronics quantum interference device (AQUID). The effective quantum dynamics of the system is proved to be that one of a two-level system. The spatial scale of the rings radii would be in the range of 5 to 20  $\mu\text{m}$ . The ring–ring interaction can be realized with the physical system of two Bose–Einstein condensates, flowing in ring-shaped optical potentials, and mutually interacting through tunnel coupling. Clearly, such systems may be relevant for quantum computation purposes, which was demonstrated further by showing how single and two qubit gates can be obtained in the setup.

The initialization of the qubit can be accomplished, for example, by imparting rotation through light-induced torque from Laguerre–Gauss (LG) beams carrying an optical angular momentum. Stacks of  $n \sim 10$  homogeneous ring lattices with tunable distance and stacks of AQUIDs have been experimentally realized (in the lab coordinated by Dumke) with spatial light modulators (SLM). Such



**Fig. 56.** Comparison between experiment and theory. (a) Comparison of the measured and calculated values of  $\Omega_0/(2\pi)$  (calculation done using GPE). The three curves correspond to calibration scales of the atomtronic SQUID radius. (b)–(d) Critical atom number as a function of the rotation rates obtained with GPE simulation and DC SQUID theory, along with the measured data and the best fit. For (b), the radius is 4.82  $\mu\text{m}$ ; for (c), the radius is 3.85  $\mu\text{m}$ ; and for (d), the radius is 2.891  $\mu\text{m}$ . Reprinted with permission from Ryu *et al.*, Nat. Commun. 11, 3338 (2020). Copyright 2020, Author(s) licensed under a Creative Commons Attribution 4.0 License.



configurations are realized by making use of the cylindrical symmetry of Laguerre–Gaussian beams and exploiting the flexibility (in terms of generating light fields of different spatial shapes) provided by the SLM devices. The stack of qubits can be realized following very similar protocols. Indeed, similar goals were carried out by realizing the AQUID with homogeneous condensates (i.e., without lattice modulation).<sup>12,15,16,67,161,604</sup> We remark that the lattice confinement brings important added values with respect to that realization. First of all, the gap between the two levels of the qubit displays a more feasible dependence with the number of atoms in the system compared with the case of homogeneous rings with a delta barrier. This is ultimately due to the fact that the barrier can be localized on a lattice spacing spatial scale;<sup>614</sup> thereby, the k-mixing that is the key feature to have a well defined two level system is not suppressed (as, in contrast, happens for homogeneous condensates with a realistic barrier. As a second positive feature, the lattice provides a platform for the qubit–qubit interaction. These two features, we believe, could ultimately facilitate the exploitation of the device in future atomtronic integrated circuits.

We also reviewed the construction of a flux qubit employing a ring condensate trapped in a regular lattice potential except for three specific lattice points with a reduced tunneling amplitude. The three weak links solution was originally suggested in quantum electronics to facilitate the function of the system as a qubit.<sup>588</sup> We apply a similar logic leading to fluxonium from the rf-SQUID: the continuous quantum fluid, in our system, is replaced by a chain of junctions connecting the different weak links. We believe that the additional lattice helps in adjusting the persistent current flowing through the system. The three weak links architecture, indeed, realizes a two-level effective dynamics in a considerably enlarged parameter space. The qubit dynamics can be read out via time-of-flight measurements. A spiral pattern emerges when the expanding atomic ring with a specific current is interfered with a reference condensate. The noise in the time-of-flight images is a hallmark of the entanglement present in the current, allowing to characterize the atomic qubit. With these methods, the type of current (entangled vs nonentangled), the magnitude and direction can be read out. The depletion at the weak-link can be used to determine the state of the qubit as well. This opens up a way to experimentally characterize atomic qubits in the lab.

Recent experiments have demonstrated the interference of currents in atomic SQUIDs for the first time. Oscillations in the critical current with applied flux are a clear hallmark of interference of atomic currents. This result has been achieved in a Bose–Einstein condensate in the dilute limit, such that it can be described within a mean-field description. While entanglement as a key ingredient for atomic qubits has not been demonstrated, this result nonetheless opens up the path to create atomic qubits via superposition of currents and observe their macroscopic entanglement.

Decoherence, of course, is an important issue for our proposal that remains to be studied. We comment, however, that measurements of the decay dynamics of a rotating condensate in an optical ring trap show that the quantized flow states have remarkably long lifetimes, of the order of tens of seconds even for high angular momentum ( $l = 10$ ).<sup>142</sup> Phase slips (the dominant mechanism of decoherence), condensate fragmentation, and collective excitations that would ultimately destroy the topologically protected quantum state are found to be strongly suppressed below a critical flow velocity. Although atom loss in the rotating condensate does not destroy the state, it can lead to

a slow decrease in the robustness of the superfluid where the occurrence of phase slips becomes more likely. We believe that the decoherence rates could be controlled within the current experimental know-how of the field. The Atomtronics’ positive trend crucially relies on the recent progress achieved in the optics microfabrication field. Thereby, central issues, of the cold atoms system, such as scalability, reconfigurability, and stability, can be feasibly addressed. In many current and envisaged investigations, there is a need to push for further miniaturization of the circuits. The current lower limit is generically imposed by the diffraction limit of the employed optics. Going to the submicrometer scale, although challenging, might be accessible in the near future. At this spatial scales, mesoscopic quantum effects could be traced out. The scalability of multiple-ring structures will be certainly fostered by tailoring optical potentials beyond the Laguerre–Gauss type (f.i. employing Bessel–Gauss laser beams). A central issue for Atomtronics integrated circuits is the minimization of the operating time on the circuit and the communication among different circuital parts (i.e., AQUID–AQUID communication). Currently, typical time rates are in the millisecond range, but a thorough analysis of the parameters controlling time rates is still missing.

XI. TRANSPORT AND DISSIPATION IN ULTRACOLD FERMION GASES

J. P. Brantut, F. Chevy, M. Lebrat, F. Scazza, S. Stringari

Atomtronics is based on the flow of quantum gases in circuits or devices. It, therefore, provides a natural framework in which transport and dissipation, two fundamental dynamical processes, can be observed, studied, and controlled. These processes are of fundamental interest in the entire field of many-body physics: first, because they involve not only equilibrium or ground state properties but chiefly that of excitations and they are intrinsically difficult to calculate from first principles. Second, for the same reason, they are very sensitive investigation tools for experimentalists. Third, they underly most of the functionalities of solid-state based quantum devices.

Fermionic quantum gases provide the most direct connection between atomtronics and solid-state electronics. The obvious analogy between the transport of fermionic atoms in light-imprinted structures and that of electrons in condensed matter systems suggests that atomtronics systems could be used as quantum simulators for their electronic counterparts, for which *ab initio* modeling is very challenging.<sup>620</sup> The vastly different scales of cold atoms, presented in Table I,

TABLE I. Comparison between cold Fermi gases and electrons in solids.

|                   | Cold Fermi gases               | Electrons in solids   |
|-------------------|--------------------------------|---|
| Interactions      | Contact, tunable               | Coulomb, with density-dependent screening                                     |
| Internal states   | Hyperfine states               | Spin  |
| Structure shaping | Light-induced                  | Gating, crystal growth  |
| Energy scales     | $E_F \sim 1 \mu\text{K}$       | $10 \text{ K} < E_F < 10^4 \text{ K}$   |
| Density scales    | $\sim 10^{12} \text{ cm}^{-3}$ | $10^{10} \text{ (semiconductors)} - 10^{22} \text{ cm}^{-3} \text{ (metals)}$ |

as well as the specific control tools make them especially promising in this perspective.

While electronic systems benefit from their ability to reach low relative temperatures, and from more than a century of development of methods and control protocols for currents and voltages, cold atomic atomtronic systems reach for previously uncharted parameter regimes: they can reach very high relative temperatures without encountering phonons or other disturbances, they offer full control and imaging of the spin degrees of freedom without the need for ferromagnetic materials and have a rich internal structure that can be leveraged as an extra “synthetic” dimension.<sup>621</sup> Of particular interest is the possibility of cold Fermi gases to operate in the strongly interacting regime, close to a Feshbach resonance, opening the possibility to realize quantum devices from strongly correlated matter bypassing the outstanding challenges encountered in solid state systems. In this regime, the system is described by the so-called BEC-BCS crossover that interpolates between weakly attractive fermions described by BCS (Bardeen–Cooper–Schrieffer) theory, and a Bose–Einstein condensate (BEC) of strongly bound molecules.<sup>622,623</sup> The equilibrium properties of gases in this regime have been extensively investigated in the last decade, and several key thermodynamic properties such as the ground state energy, critical temperature, or pairing gap are now known with high precision.<sup>624,625</sup>

The recent years have seen a growing number of experiments exploring the dynamics of fermionic gases in this strongly interacting regime. New systems mixing bosonic and fermionic superfluids provides renewed opportunities to study superfluid flow.<sup>626–628</sup> The development of two-terminal systems for cold atoms in particular provides the simplest device-like geometry.<sup>498</sup> For a long, low dimensional channel, this has allowed for the measurement of particle,<sup>629</sup> spin,<sup>630</sup> and heat conductances<sup>631</sup> as well as off-diagonal transport coefficients such as spin-drag or thermopower. For short, planar junctions, it realizes a tunnel connection for which superfluidity yields the celebrated Josephson effect.<sup>33,430,437,438</sup>

This contribution presents some of the most recent development of ultracold Fermi gases in the atomtronics context. In Sec. XI A, transport phenomena in superfluid Fermi gases are discussed, first in the perspective of the Landau criteria, then in the case of Josephson junctions. In Sec. XI B, we describe transport of Fermi gases in mesoscopic channels. In Sec. XI C, the physics of the fast spin drag in normal Fermi gases is presented.

## A. Superfluid transport with Fermi gases

### 1. Fermionic superfluidity and critical velocity

The first microscopic theory of dissipation in superfluids was proposed by Landau who predicted the existence of critical velocity below which an object in motion in a superfluid feels no drag.<sup>632</sup> Landau’s original argument was based on constraints imposed by energy and momentum conservation when elementary excitations are shed in the superfluid. In this limit, the critical velocity is given by the minimum value of  $\omega(k)/k$ , where  $\omega(k)$  is the dispersion relation of low-energy modes of the system. For a concave dispersion relation, this is the slope of  $\omega(k)$  at the origin and the critical velocity is, therefore, simply the sound velocity.

In fermionic systems, the excitation spectrum is composed of both bosonic collective modes (the so-called Bogoliubov–Anderson

modes) corresponding to phonons and fermionic quasi-particles associated with broken Cooper pairs.<sup>633</sup> These two sectors lead to different predictions for the critical velocity when interactions are varied across the BEC-BCS crossover. In the BEC regime, where pairs are tightly bound, phonons set the critical velocity, as in a traditional atomic Bose–Einstein condensate. On the contrary, on the BCS side of the resonance, the Cooper pairs are loosely bound and are easily broken by a moving object. In this regime, the critical velocity is  $v_c \simeq \Delta/p_F$ , where  $\Delta$  is the excitation gap and  $p_F$  is the Fermi momentum. The existence of these two excitation branches leads to a maximum of Landau critical velocity close to the unitary limit that was experimentally observed by stirring an optical potential in the cloud.<sup>634,636</sup>

Recent experiments on atomic mixtures have raised the question of the onset of dissipation in two counterflowing superfluids.<sup>626,628,636</sup> Experiments on dual Bose–Fermi superfluids revealed the existence of a critical velocity which was later on interpreted as an extension of Landau’s seminal argument similar to parametric down-conversion in quantum optics. In this scenario, the relative motion of the two superfluids can excite *pairs* of excitations in the superfluids.<sup>627,637–639</sup> This modifies the expression of the critical velocity, which is equal to the sum of the sound velocities of the two superfluids when phonons limit superfluidity, a prediction that agrees with experimental measurements<sup>628</sup> performed on mixtures of <sup>6</sup>Li and <sup>7</sup>Li.

Let us conclude this subsection by stressing some of the hypotheses underlying Landau’s scenario. First, as mentioned earlier, the identification of Landau critical velocity in the phonon sector with sound velocity assumes that the dispersion relation is convex. Although this is true for bosons in free space, this is no longer the case for fermions, for which the coupling with the broken-pair particle–hole continuum bends the dispersion relation downwards.<sup>640</sup> Likewise, the presence of a transverse trapping in experiments leads to a reduction of the critical velocity due to an inversion of the concavity of the dispersion relation at large momenta, a feature first pointed out in weakly interacting Bose–Einstein condensates<sup>641,642</sup> and recently generalized to arbitrary hydrodynamic superfluids.<sup>643</sup> Second, Landau’s argument assumes that the velocity of the moving disturbance is constant, while in experiments, the motion of the disturbing potential is usually oscillatory to account for the finite size of the system. By analogy with an accelerated electric charge that radiates electromagnetic wave at an arbitrary small velocity, Landau critical velocity is suppressed for accelerated disturbances.<sup>644</sup> Finally, as initially proposed by Feynman and Onsager,<sup>645,646</sup> topological defects, such as quantized vortices, are responsible for the onset of dissipation for stronger disturbances.<sup>495</sup>

### 2. Josephson currents

The Josephson effect represents a quintessential manifestation of macroscopic quantum phase coherence, stemming from spontaneous symmetry breaking in superfluid states. A so-called Josephson junction is typically created by weakly coupling two superfluid order parameters through a thin insulating barrier. In the solid state, this is achieved by separating two superconductors with a nanometer-sized insulating layer. Josephson first predicted that a dissipationless supercurrent  $I_s = I_c \sin(\varphi)$  should flow across a tunnel junction in the absence of an applied voltage, associated with the coherent tunneling of Cooper pairs and sustained only by the relative phase  $\varphi$  between the two order parameters. The maximum value  $I_c$  of the supercurrent is coined by

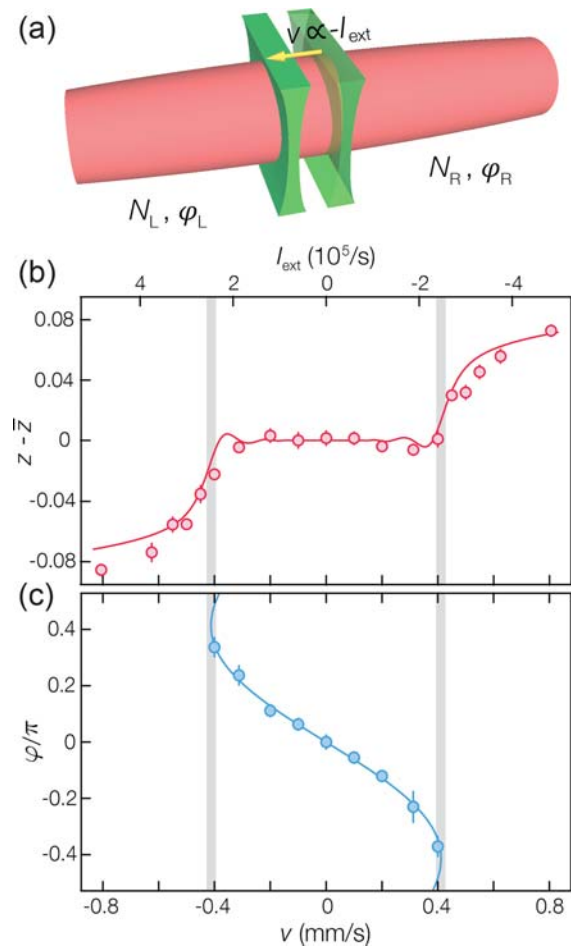
the Josephson critical current, and it is directly related to the strength of the tunnel coupling between the two order parameters within the insulating barrier. The measurement of  $I_c$  provides a powerful probing tool offering fundamental insights into the microscopic properties of the involved superfluid states and their robustness against dissipation. For example, for BCS superconductors,  $I_c$  is directly related to the order-parameter amplitude, i.e., the gap  $|\Delta|$ , by the Ambegaokar-Baratoff relation. For applied currents above  $I_c$ , the junction enters a resistive regime, where a nonzero stationary conductance arises from dissipative excitation processes and a finite electro-chemical potential response is generated across the junction.

Experimental studies with atomic superfluids have so far mainly targeted coherent transport in BECs with various geometries and optically engineered weak links.<sup>12,16,67,391,428,448,604,647</sup> On the other hand, the study of supercurrents between weakly coupled superfluid Fermi gases is of high relevance from both the fundamental and the practical point of view,<sup>430,648,649</sup> since transport therein is crucially influenced and complicated by strong interparticle interactions and their interplay with fermionic statistics. Only recently, dc Josephson supercurrents have been observed in strongly interacting Fermi gases close to a Feshbach resonance (Fig. 57).<sup>33</sup> Reminiscent of the behavior of the Landau critical velocity across the BCS-BEC crossover,<sup>622</sup> the Josephson critical current was found to exhibit a pronounced maximum around unitarity, resulting from opposite variations of the chemical potential and the pair condensate fraction, the latter playing the role of the order-parameter amplitude throughout the crossover.<sup>33</sup> First experimental investigations of the Josephson effect in quasi-two-dimensional fermionic condensates have also been reported recently, providing information on the connection between condensation and the Berezinskii-Kosterlitz-Thouless superfluid transition.<sup>96</sup> Moreover, experiments showed the breakdown of coherent Josephson transport to be accompanied by the nucleation of topological defects generated above critical flows by the barrier constriction and subsequently emitted into the superfluid bulk.<sup>437,438</sup> More efforts will be necessary to shed light on the precise mechanisms underlying dissipation in Josephson junctions between crossover Fermi superfluids, and on the interplay of bosonic and fermionic excitation mechanisms, corresponding to the Bogoliubov-Anderson and pair-breaking excitation branches observed for an obstacle moving through the superfluid.<sup>495,633-635</sup> Such explorations will be essential for our understanding of dissipative transport in highly correlated fermionic systems and for extending the applications of the Josephson effect to emerging atomtronic devices.

## B. Fermionic transport in mesoscopic channels

Mesoscopic devices refer to small-size systems whose transport properties are influenced by the quantum nature of matter. For example, the conductance of a narrow constriction becomes quantized when its width is comparable to the de Broglie wavelength of the particles traversing it. Initially demonstrated with electrons in semiconducting nanostructures,<sup>650</sup> mesoscopic transport can be naturally extended to fermionic atoms.

As quantum gases have to be particularly shielded from environmental perturbations, they are intrinsically closed systems, which is both a blessing and a complication to study mesoscopic transport phenomena. On the one hand, the relaxation of thermodynamical quantities involved in transport such as momentum or spin mostly depend on interparticle interactions, which can be tuned for instance via



**Fig. 57.** Realization of a current-biased Josephson junction between ultracold fermionic superfluids. (a) Two superfluid reservoirs (L, left; R, right) of  $^6\text{Li}$  fermion pairs are weakly coupled through a thin optical repulsive barrier created using a DMD. An external current  $I_{\text{ext}}$  is imposed by translating the tunneling barrier at a constant velocity  $v$ . Pair transport is tracked by recording the relative imbalance  $z = (N_R - N_L) / (N_R + N_L)$  through *in-situ* absorption imaging, while the order-parameter relative phase  $\phi$  is revealed through matter-wave interference. (b) Experimental current-imbalance characteristic, and (c) current-phase relation  $I(\phi)$  for a crossover Fermi gas on the BEC side of the Feshbach resonance. The solid line denotes the fit to a resistively-shunted Josephson junction circuit model, while the shaded vertical lines indicate the extracted  $I_c$ . Reprinted with permission from Kwon *et al.*, *Science* **369**, 84 (2020). Copyright 2020, AAAS.

Feshbach resonances. On the other hand, real-life transport measurements with electrons imply connecting macroscopic leads acting as particle and heat baths to a smaller system of interest, usually treated as a grand canonical ensemble. With quantum gases, such a paradigm for transport requires partitioning the isolated system into a mesoscopic conductor and two or more macroscopic reservoirs that thermalize fast enough compared to the transport timescales to be considered in thermodynamical equilibrium. Cold-atom realizations close to such multi-terminal setups include single and multiple Josephson junction arrays of trapped BECs,<sup>448,651</sup> weak links in ring traps,<sup>16,161</sup> and planar junctions between two fermionic superfluids.<sup>33,430,437</sup>



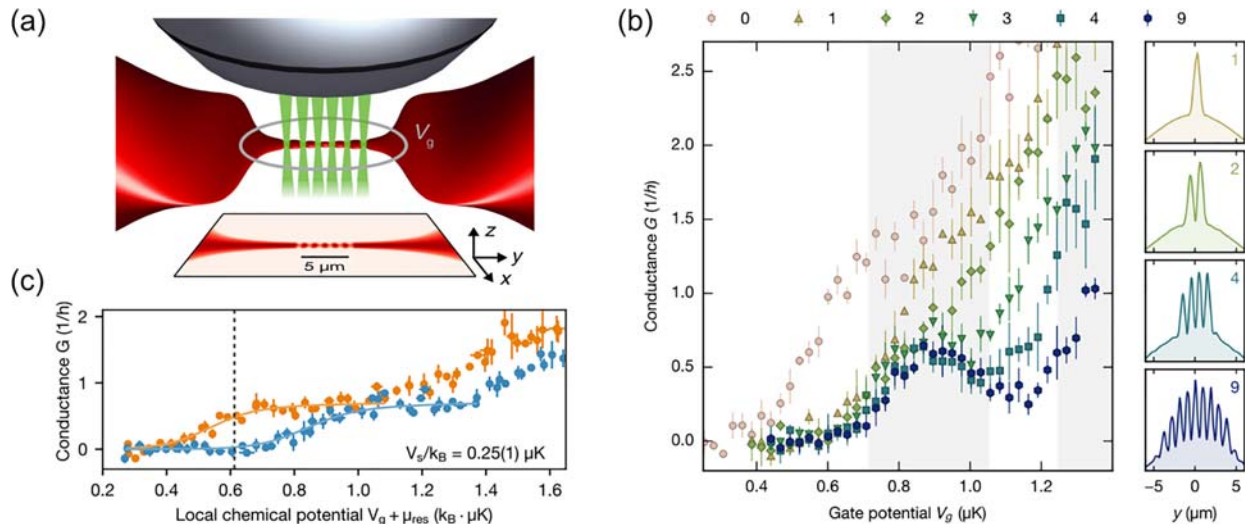
By reducing the dimensionality of the mesoscopic region, the atomic equivalent of quantum point contacts has been realized, displaying quantized conductance.<sup>652</sup> Starting from this two-terminal configuration, more complex structures can be engineered by projecting arbitrary light potentials via holographic techniques.<sup>95</sup> Recently, this technique allowed to investigate the insulating properties of a mesoscopic lattice imprinted site by site within a quantum wire,<sup>653</sup> visible as a suppression of conductance at Fermi energies located in the lattice band gap [Figs. 58(a) and 58(b)]. Strikingly, this insulating behavior persists as attractive interactions are increased to the point where reservoirs become superfluid. The robustness of the fermionic character of transport can be attributed to the existence of a Luther–Emery liquid,<sup>654</sup> a strongly correlated phase of matter distinctive of the 1D character of the channel. In a more recent set of experiments, optical control in atomic point contacts was extended to spin by using light tuned close to atomic resonance to create local effective Zeeman shifts [Fig. 58(c)]. This leads to the realization of an ideal spin filter with cold atoms, one of the most fundamental spintronic devices.<sup>655</sup> In the presence of weak interactions, near-resonant light scattering can be entirely accounted for by including losses in a Landauer–Büttiker model.<sup>656</sup> Such a progress toward spin-dependent transport opens avenues for exploring the transport dynamics of strongly correlated heterostructures, where novel nonequilibrium spin and heat transport<sup>657,658</sup> and exotic phases of matter<sup>659</sup> could be observed.

### C. Fast spin drag in normal Fermi gases

Spin drag is a ubiquitous concept in many branches of physics. It is usually associated with interaction effects that affect the Euler equation for the spin current. Spin drag can be of collisional nature, giving

rise to spin diffusion since collisions do not conserve the spin current, or of collisionless nature, being at the origin of nondissipative dynamics.<sup>660–667</sup> Experiments on transverse spin diffusion<sup>663</sup> (Leggett–Rice effect) in an ultracold Fermi gas along the BCS–BEC crossover have allowed for the determination of the relevant combination  $G_0 - G_1/3$  of the spin parameters of Landau theory of Fermi liquids. In particular, the parameter  $G_1$  accounts for the strength of spin–current interactions. Spin drag can also be due to the modification of the equation of continuity in the spin channel caused by interactions and yielding a violation of the corresponding f-sum rule. An example of such a spin drag behavior of collisionless nature (hereafter called “fast spin drag”) takes place in the Andreev–Bashkin effect caused by quantum fluctuations in a mixture of two interacting superfluids.<sup>668,669</sup> This effect is very tiny and difficult to observe in dilute Bose gases, unless one considers one-dimensional configurations<sup>670</sup> or quantum gases in an optical lattice.<sup>669</sup> In the following, we will discuss some consequences of fast spin drag in a normal (nonsuperfluid) interacting mixture of two Fermi gases, where the effect can be sizable and hopefully measurable. Other nontrivial examples of fast spin drag concern the dynamical behavior of coherently coupled Bose–Einstein condensed mixtures.

To investigate the phenomenon of fast spin drag, it is convenient to consider an external perturbation of the form  $H_{\text{pert}} = -\lambda f(\mathbf{r}) \Theta(t)$  applied to the system, where  $\Theta(t)$  is the usual Heaviside step function (equal to 0 for  $t < 0$  and 1 for  $t > 0$ ) and the function  $f(\mathbf{r})$  characterizes the nature of the perturbation, while  $\lambda$  is its strength. For example, in an ultracold atomic gas, a convenient choice is  $f(\mathbf{r}) = x$ , corresponding to a boost generated by an optical potential. If the perturbation is equally applied to both components of the mixture, the velocity acquired by the system is given, for short times, by  $v_x = \lambda t/m$ , where



**FIG. 58.** Studying mesoscopic transport with ultracold fermions. (a) A degenerate Fermi gas of lithium-6 atoms is shaped using repulsive light potentials into a one-dimensional channel smoothly connected to two macroscopic reservoirs. Spatial light modulation techniques combined with high-resolution optics allow to imprint additional structures to the 1D channel, such as a lattice of variable length. (b) For weak interactions, the current through the lattice is proportional to applied bias and particle transport is captured by a linear conductance coefficient. Conductance shows a local minimum as a function of the overall chemical potential, indicating the emergence of a band gap when approaching the infinite lattice limit. (c) Spin-dependent quantized conductance in the presence of a near-resonant obstacle focused on the 1D channel, realizing the cold-atom equivalent of a spin filter. Panels (a) and (b) reprinted with permission from Lebrat *et al.*, Phys. Rev. X **8**, 011053 (2018). Copyright 2018, American Physical Society. Panel (c) reprinted with permission from Lebrat *et al.*, Phys. Rev. Lett. **123**, 193605 (2019). Copyright 2019, American Physical Society.



$m$  is the atomic mass, and we have set  $\hbar = 1$ . A more interesting scenario occurs when the perturbation is applied in a selective way only to one component (hereafter called component 1). In this case, the velocities acquired by the two components will be different, and for short times, they can be easily calculated starting from the many-body wave function of the system which, in the presence of a fast perturbation, takes the form  $\Psi(t) = \exp[i\lambda t x_1 - iHt]\Psi_0$  with  $x_1$  being the center-of-mass operator relative to the first component. The velocities acquired by the two components are then given by

$$v_1 = \frac{d}{dt} \langle x_1 \rangle = 2\lambda t \langle [x_1, [H, x_1]] \rangle / N = \lambda t (m_1^n + m_1^s) / N, \quad (99)$$

$$v_2 = \frac{d}{dt} \langle x_2 \rangle = 2\lambda t \langle [x_1, [H, x_2]] \rangle / N = \lambda t (m_1^n - m_1^s) / N, \quad (100)$$

where  $N$  is the total number of particles and we have expressed the double commutators as proper combinations of the energy-weighted moments  $m_1^n = \int d\omega \omega S(x_1 + x_2, \omega)$  and  $m_1^s = \int d\omega \omega S(x_1 - x_2, \omega)$  of the dynamic structure factors relative, respectively, to the *in-phase* ( $x_1 + x_2$ ) and *out-of-phase* ( $x_1 - x_2$ ) operators.

In the absence of current interactions, only component 1 will feel the external kick. In fact, in this case, the commutator  $[H, x_2]$  commutes with  $x_1$  being uniquely fixed by the kinetic energy term in the Hamiltonian and, consequently,  $v_2 = 0$  and  $m_1^n = m_1^s$ . While the *in-phase* energy-weighted moment  $m_1^n$  is given by the model-independent f-sum rule  $m_1^n = N/2m$ , as a consequence of Galilean invariance, the *out-of-phase* moment is instead sensitive to the presence of spin-current interactions,<sup>671</sup> and in uniform matter, it takes the value

$$m_1^s = N \frac{1}{2m^*} (1 + G_1/3), \quad (101)$$

where  $m^* = m(1 + F_1/3)$  is the effective mass of quasi-particles fixed by the Landau's parameter  $F_1$  accounting for density-current interactions and determining the low-temperature behavior of the specific heat,<sup>672</sup> while  $G_1$  is the spin-current interaction parameter. Result [Eq. (101)] reflects the fact that particle-hole excitations, properly accounted for by Landau's theory of Fermi liquids, do not exhaust the f-sum rule, multipair excitations playing a crucial role in providing the remaining difference. The spin energy-weighted sum rule [Eq. (101)] was actually employed<sup>673</sup> to estimate the average multipair excitation energy in liquid <sup>3</sup>He.

The violation of the f-sum rule in the spin channel is responsible for the spin drag effect in a normal Fermi liquid, which, according to Eqs. (99)–(100), takes the form

$$\frac{v_2}{v_1} = \frac{(F_1 - G_1)/3}{2 + (F_1 + G_1)/3}, \quad (102)$$

revealing that the drag effect ( $v_2 \neq 0$ ) vanishes only if the Landau parameters  $F_1$  and  $G_1$  are equal. In a dilute Fermi gas the values of the Landau's parameters are available in both three<sup>674,675</sup> and two dimensions,<sup>676</sup> using second-order perturbation theory. For example, in 3D, one has

$$F_1 = \frac{8}{5\pi^2} (7 \ln 2 - 1) (k_F a)^2; \quad G_1 = -\frac{8}{5\pi^2} (2 + \ln 2) (k_F a)^2, \quad (103)$$

showing explicitly that fast spin drag is quadratic in the dimensionless parameter  $k_F a$ , where  $a$  is the *s*-wave scattering length and  $k_F$  is the

Fermi wave vector. Similarly to the Andreev–Bashkin effect in interacting superfluids,<sup>668,669</sup> the fast spin drag exhibited by a normal Fermi gas has consequently a typical beyond-mean-field nature.

The applicability of Landau theory of Fermi liquids, yielding result (102) for the fast spin drag, holds for temperatures much smaller than the Fermi temperature  $T_F$ . At the same time, the temperatures should be higher than the critical temperature for superfluidity. These conditions can be well experimentally satisfied in the BCS regime of small and negative scattering lengths. For larger values of  $k_F |a|$ , when the system approaches the unitary regime, its applicability is instead questionable because the superfluid critical temperature is of the order of the Fermi temperature. The experimental determination of the fast spin drag effect along the BEC-BCS crossover would then complement the measurements of the Leggett–Rice effect,<sup>663</sup> providing a crucial test of Landau's theory and stimulating further theoretical work on spin transport phenomena.

## D. Concluding remarks and outlook

The investigation of transport and dissipation in Fermi gases has only started recently, and many new directions are already emerging. The available light-shaping techniques allow, in principle, for complex geometries to be investigated.<sup>7,59</sup> A particularly appealing situation is the ring trap, which has been successfully explored for weakly interacting bosons.<sup>390</sup> Complex geometries are accessible using the concept of synthetic dimensions,<sup>677</sup> where multiterminal geometries are naturally arising from two physical terminals.<sup>621</sup> Transport of correlated fermions in optical lattices has started recently in bulk lattice systems with promising results on the quantum simulation of the Hubbard model.<sup>666,678,679</sup>

The intrinsically low energy scales also implies that currents are weak: translated into electronic scales, the typical currents of fermionic particles in a single mode conductor amounts to fractions of femto-amperes. Reaching a signal-to-noise ratio comparable with that achieved in the condensed matter context, which would allow for a direct validation of a quantum simulation approach to transport, calls thus for new methods of detection. A practical route is the combination of existing transport systems with single atom sensitive methods that have been demonstrated already, such as fluorescence based counting<sup>680–682</sup> or quantum-gas microscopy.<sup>683</sup> These methods provide ultimate sensitivity, but still suffer from the technical effects of sample-to-sample preparation noise that scales unfavorably with the number of particles. Ultimately, the limit to signal-to-noise is set by measurement back-action. Achieving this limit would then allow for the reconstruction of the full counting statistics of transport process. Several schemes interfacing atoms with photons in a cavity allow, in principle, to achieve this limit either in the lattice context<sup>684</sup> or for the two-terminal configuration,<sup>685</sup> and experiments directed at implementing such methods have already started.

The physics of complex atomtronics devices featuring Fermi gases with strong interactions opens many possibilities, also of interest in the condensed matter community at large. An overarching goal is the manipulation of topological superfluids,<sup>686</sup> such as *p*-wave superfluids or Kitaev chains,<sup>687</sup> which would provide an avenue to study topological protection of quantum information, thus guiding the development of topological quantum computers.

## XII. TRANSPORT IN BOSONIC CIRCUITS

**T. Haug, R. Dumke, L.-C. Kwek, W. von Klitzing, L. Amico**

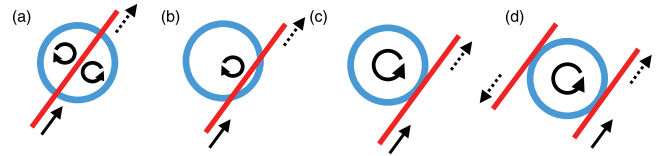
Atomtronics opens up a new approach to study fundamental problems of transport of quantum matter in various settings with widely different light-controlled atomic circuits.<sup>4</sup> Of particular interest is transport generated by attaching a circuit to reservoirs that induce a directed current through the system. Transport between atomic reservoirs has been studied to realize fundamental condensed matter systems.<sup>498,629,649,652</sup> Furthermore, precise control of the light potentials allows one to transport bosonic fluids at hypersonic speeds in ring circuits<sup>117,128</sup> and in a coherence preserving manner.<sup>688</sup> Often these basic atomic circuits can be understood by using a simplified lumped element model that relies only on a few elements.<sup>29</sup> From there, larger circuits composed of many basic circuits could be constructed to realize large scale atomtronic networks. To this end, there is a considerable interest in understanding the transport through basic circuit elements.

Recent studies investigated the transport and dynamics in other circuits like rings and Y-junctions.<sup>118,689–693</sup> These systems have been well studied in electronic setups. Surprising differences arise with bosonic atoms instead of fermionic electrons: Andreev reflections, known from superconductor–metal interfaces, can also occur at the interface of two bosonic condensates: If the density wave excitation in a one-dimensional condensate is transmitted from the first to second condensate, a hole (an excitation with negative amplitude) is reflected back into the first condensate.<sup>690,694–697</sup> For ring circuits, Aharonov–Bohm oscillations occur in the current for electronic systems when an a magnetic field is applied to the ring.<sup>698</sup> For bosonic rings, this Aharonov–Bohm effect does not occur.<sup>700,715</sup> As a first step to observe this effect in cold atoms, a recent experiment demonstrated nonreciprocal transport mediated by artificial magnetic fields in closed loops.<sup>699</sup> Transport can also be achieved by driving the system parameter in time. Here, topological pumping has been shown to be a robust way to generate transport.<sup>702,719</sup> Here, the circuit parameters are driven periodically in time such that a directed transport arises, which is protected by topological features of the system.<sup>702–704</sup> For ring systems with applied flux, the transmission becomes fractional in atom number, and highly entangled states can be generated in the process.<sup>691</sup> This promises important applications in quantum-enhanced sensing and quantum information.

First, we discuss recent advances in matter-waveguides, which allow one to transport cold atoms over long distances (see Sec. XII and Fig. 59). Then, we review the transport in two elemental Atomtronic circuits (see Fig. 60): A ring attached to leads (see Figs. 61–63) and a Y-junction (see Figs. 64 and 65). We investigate different limits: Atoms prepared in a nonequilibrium initial state with all atoms on one side of the system. Second, density wave excitations that propagate through a system filled with atoms. Finally, we also shortly mention topological pumping of atoms in atomtronic circuits (see Fig. 66).

### A. Matterwave guides

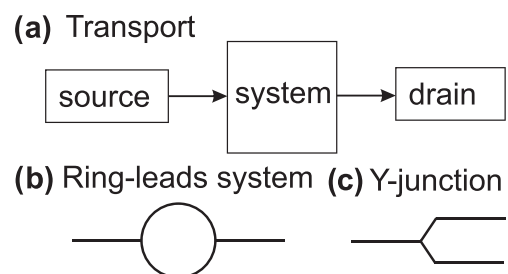
A perfect waveguide allows the guided wave to travel undisturbed over any distance. In practice, there are always imperfections such as absorption and spatial variations of the guiding potential. For matter-wave guides based on electro-magnetic potentials, absorption plays no



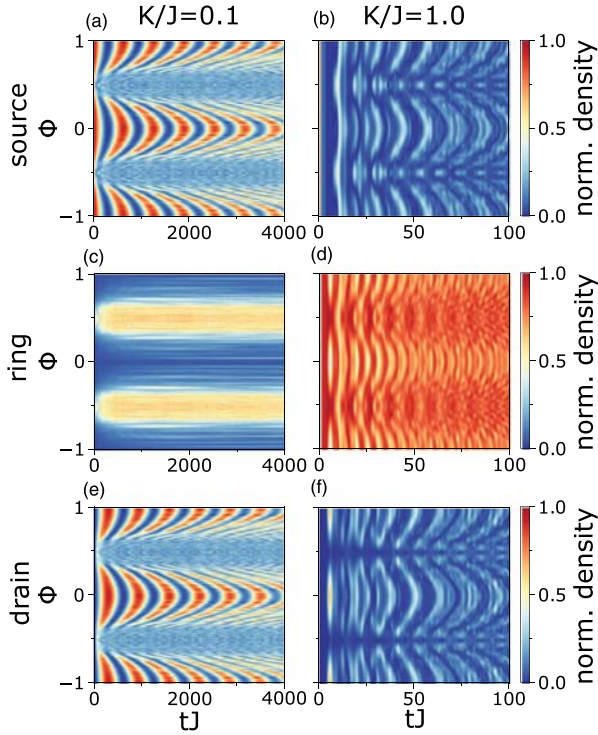
**Fig. 59.** Possible configurations for coupling of a TAAP ring (blue) to an optical guide potential (red). In all cases the atoms are in a magnetically or rotational sensitive state in the TAAP potential, which is then coupled to a magnetically or rotational nonselective state by tunnel coupling or a spatially selective microwave transition. (a) symmetric ring-lead system, (b) asymmetric ring-lead system. c) tangential configuration, where dipole is coupled to only one direction of rotation in the ring. (d) TAAP ring coupler between two dipole wave guides. Note that in (a) and (b) the coupling on the two sides of the ring can be individually tuned.

role. In most cases, the shapes of the guiding potentials are defined either by a physical structure such as wires in the case of magnetic potentials or by light-fields in the case of dipole traps. Examples include, imperfections in the shapes of the wire,<sup>106</sup> the grain size of the copper<sup>705</sup> and for the dipole finite amplitude control,<sup>32</sup> diffraction and speckles. There are a number of solutions to reduce the impact of the imperfections of magnetic waveguides, such as improved manufacturing techniques and periodic current reversal.<sup>106</sup> Optical techniques employ feedback to image the potential using cold atoms and then correct the imperfections in a feedback loop.<sup>32</sup> Nevertheless, since some level of imperfection in the magnetic wire structure or in the dipole imaging system is unavoidable, waveguides created by artificial structures will always have a certain degree of roughness.

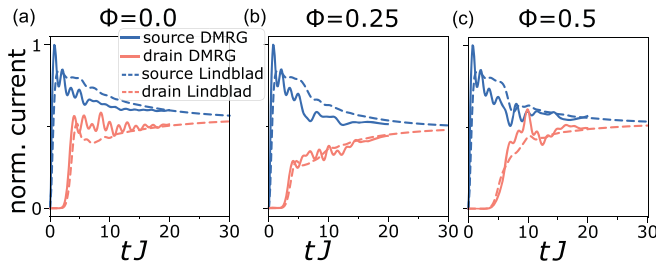
The effect of these imperfections is characterized by the spatial wavelength  $\lambda$  and amplitude  $a(k)$  of the modulation of the waveguide potential: A tighter bend will have a stronger effect than a very smooth one. For optical traps, this strength can be calculated directly by estimating the level of control one has over the optical potential, e.g., by imaging the speckles or by estimating the noise level in the feedback to the shape of the waveguide. In the case of magnetic waveguides, the variation of the potential can be imaging the break-up of a Bose–Einstein condensate, which is brought close to the wires. An absolute scale can be established from the resulting images via chemical potential of the BEC.



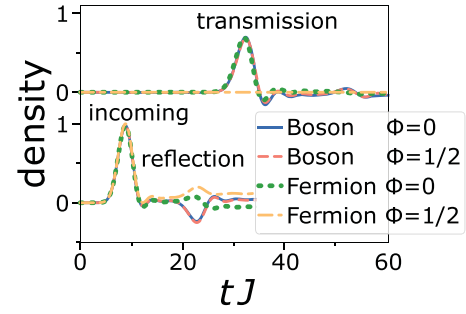
**Fig. 60.** (a) General transport setup, composed of a source, system and drain. Atoms flow from source, via the system to drain. The current flowing through the system is the quantity of interest, that reveals fundamental features of the system. Specific examples of this kind of setup studied here are a (b) ring-leads system or a (c) Y-junction.



**Fig. 61.** Time evolution of density in source (a) and (b), ring (c) and (d), and drain (e) and (f) plotted against flux  $\Phi$ . (a), (c), and (e) weak ring-lead coupling  $K/J = 0.1$  (on-site interaction  $U/J = 5$ ). (b), (d), and (f) strong ring-lead coupling  $K/J = 1$  ( $U/J = 0.2$ ). Time is indicated  $tJ$  in units of inter-ring tunneling parameter  $J$ . The number of ring sites is  $L = 14$  with  $N_p = 4$  particles initially in the source. The density in the ring is  $n_{\text{ring}} = 1 - n_{\text{source}} - n_{\text{drain}}$ . Reprinted with permission from Haug *et al.*, Phys. Rev. A **100**, 041601 (2019). Copyright 2019, American Physical Society.



**Fig. 62.** Current through the Aharonov-Bohm ring. (a)–(c) Evolution of source and drain current toward the steady state with DMRG (solid line) and Lindblad formalism (dashed) for hard-core bosons,  $K = 1$  and  $L_R = 10$ . For DMRG, both reservoirs and ring are solved with Schrödinger equation as a closed system. Source and drain are modeled as chains of hard-core bosons with equal length  $L_S = L_D = 30$ . Initially, the source is prepared at half-filling ( $N_p = 15$ ) in its ground state (ring and drain are empty) decoupled from the ring [ $K(t = 0) = 0$ ]. For  $t > 0$ , the coupling is suddenly switched on [ $K(t > 0) = J$ ]. This setting creates highly nonequilibrium dynamics. Due to numerical limitations, we analyze the short-time dynamics. For the open system, the reservoirs obey Pauli-principle with  $r = 0.65$  and  $\Gamma = 1.5$ . Reprinted with permission from Haug *et al.*, Quantum Sci. Technol. **4**, 045001 (2019). Copyright 2019, IOP Publishing Ltd. Reprinted with permission from Haug *et al.*, Phys. Rev. A **100**, 041601 (2019). Copyright 2019, American Physical Society.



**Fig. 63.** Propagation of a small density excitation in a ring-lead system for hard-core bosons and spinless fermions for zero and half-flux. Fermion transmission is flux-dependent, while hard-core bosons are flux independent. The source and drain lead has length  $L_S = L_D = 80$  and the ring  $L_R = 40$ , the particle number  $N = 100$ , strong coupling with  $K = J$  and  $\varepsilon_D = 0.3J$ . The propagation of the density excitation in time. The upper curves show the transmitted density wave into the drain lead (integrated between site 130 and 135), and the lower curves the incoming and reflected wave in the source lead (65 and 70). The background density is subtracted. Curves stay constant in area shadowed by legend. Reprinted with permission from Haug *et al.*, Quantum Sci. Technol. **4**, 045001 (2019). Copyright 2019, IOP Publishing Ltd.

Increasing the distance between the atoms and the current-carrying conductor and decreasing the transverse trapping frequency reduce the roughness of the waveguide. For distances ( $d$ ) from the waveguide much larger than the characteristic wavelength ( $\lambda$ ), this reduction ( $K$ ) in roughness can be estimated as a function of the spatial frequency ( $k = 2\pi/\lambda$ ) as<sup>705</sup>

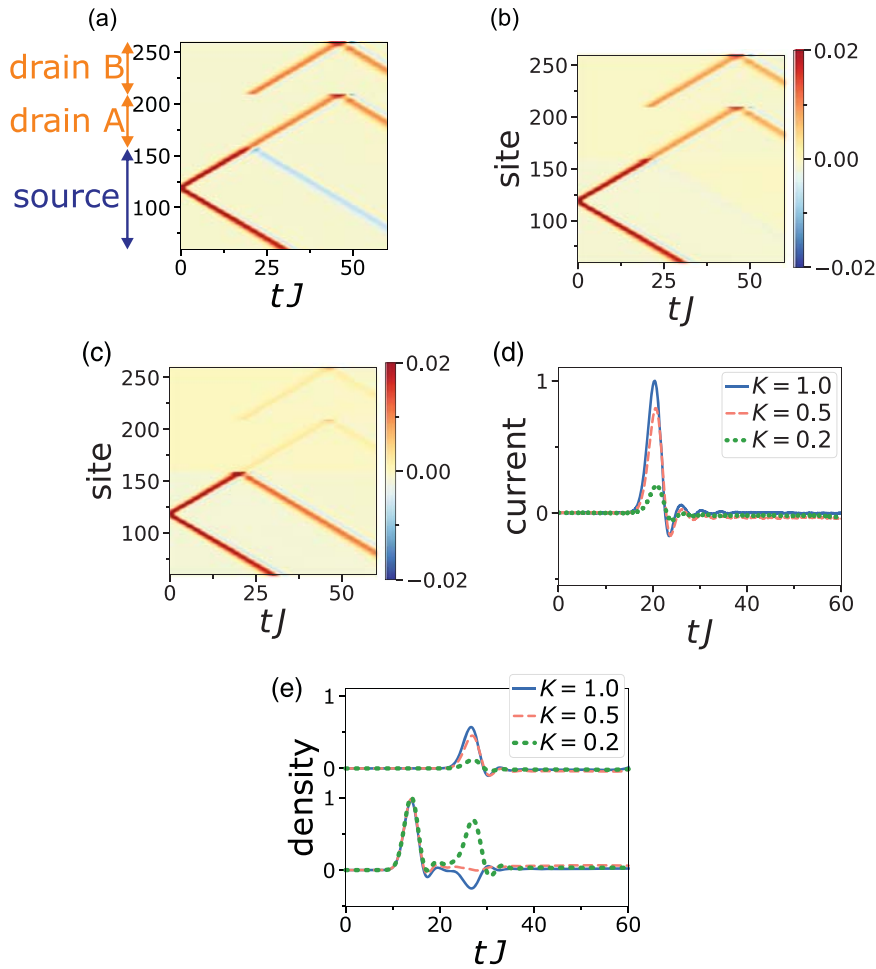
$$K(d, \lambda) = (kd)^{-1/2} e^{-kd}. \quad (104)$$

By increasing  $d$ , it was possible to observe interference fringes between two condensates after propagating them on a magnetic atomchip waveguide for up to  $120 \mu\text{m}$ , albeit reducing the transverse trapping frequency from the kilohertz level down to  $\omega_{\perp} = 2\pi \cdot 120 \text{ Hz}$ .<sup>706</sup> If no propagation is required, even spatially modulated waveguides can exhibit robust coherence.<sup>183</sup>

Very smooth optical dipole matterwave guides can be achieved by weakly focusing a laser beam and taking care to avoid laser speckles. If the imaging system is Fourier limited, then it cannot produce any structure smaller than the focus, resulting in a perfectly smooth waveguide. By the same token, however, no structure other than a simple linear waveguide can be produced by this method.

A different approach has been recently demonstrated, where the shape of a ring-shaped waveguide is defined by modifying a simple DC quadrupole field using only homogeneous audio-frequency and radio-frequency fields.<sup>117</sup> In the so-called time-averaged adiabatic potentials, the radial and vertical confinement is limited to a ring and the maximum spatial azimuthal frequency that can be addressed is  $\phi = 4\pi$ . Since the field generating magnets only have to produce homogeneous and quadrupole fields, they can be far away. Equation (104) predicts a reduction of the field modulations down to a factor  $10^{-138}$  of their strength at the magnets, thus practically eliminating them. This has made it possible to propagate Bose-Einstein condensates over distances of more than 10 cm without causing any additional heating.

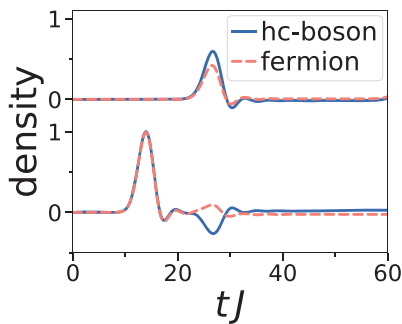




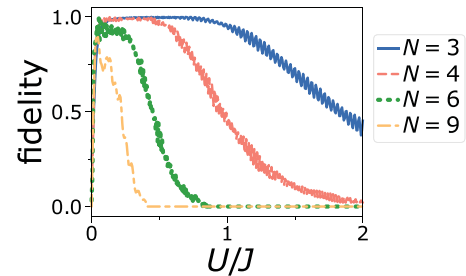
**FIG. 64.** Propagation of small excitation in a Y-junction for the hard-core boson model. The source lead has length  $L_S = 160$ , the drain lead each  $L_D = 50$ , the particle number  $N = 130$ , initial distance of the excitation to the junction  $d = 40$  and  $\varepsilon_D = 0.3J$ . The source lead is from site 1 to 160, the first drain lead from 160 to 210, and the second one from site 210 to 260. The coupling at the junction (site 160) is (a)  $K = 1J$ , (b)  $K = 0.5J$ , (c)  $K = 0.2J$ . (d) Current through the junction [Eq. (108)] in time. (e) The propagation of the density excitation in time. The upper curves show the transmitted density wave into the drain lead (integrated between site 170 and 175), and the lower curves the incoming and reflected wave in the source lead (145 and 150). The background density is subtracted. For  $K = 1J$  (solid) we observe a negative reflection (Andreev-like),  $K = 0.5J$  (dashed) nearly no reflection,  $K = 0.2J$  (dots) a large positive reflection amplitude. Reprinted with permission from Haug *et al.*, Quantum Sci. Technol. **4**, 045001 (2019). Copyright 2019, IOP Publishing Ltd.

A very interesting perspective is to combine the TAAP rings with optical potentials. The standard way to load atoms into the TAAP ring is currently to transfer them from an optical dipole trap.<sup>117,118</sup> Using radio-frequency or microwaves, it would be possible to create a beam

splitter between the ring and the optical potential. This couples the magnetically or rotational sensitive state in the ring to a magnetically and rotational nonsensitive one in the optical guide. Possible configurations would include, for example, a (a)-symmetric ring-lead system



**FIG. 65.** Comparison between hard-core bosons and spinless fermions for the transmission and reflection of a small density excitation in a Y-junction. Bosons show clear negative Andreev-reflection (solid and dashed curve in center bottom), in contrast to fermions. The source lead has length  $L_S = 160$ , the drain lead each  $L_D = 50$ , the particle number  $N = 130$ ,  $d = 40$ ,  $K = J$ , initial half-filling and  $\varepsilon_D = 0.3J$ . The propagation of the density excitation in time. The upper curves



**FIG. 66.** Numerical simulation of fidelity of creating a NOON-like entangled state by pumping  $N$  particles through a simplified ring-lead system of three lattice sites (see the text).  $N$  particles are placed initially in the source lead, then pumping is switched on, which transport particles into the ring. Then, fidelity of creating a NOON state (particles are in an entangled state of being in either of the two paths of the ring). Fidelity is plotted against interaction  $U$  in units of intersite hopping  $J$ . Reprinted with permission from Haug *et al.*, Commun. Phys. **2**, 127 (2019). Copyright 2019, Author(s) licensed under a Creative Commons Attribution 4.0 License.

[Figs. 59(a) and 59(b)], a dipole guide coupled tangentially to the ring [Fig. 59(c)]. Since the diameter of TAAP rings and, therefore, their resonant angular momentum is easily tuned, they could act as a velocity-selective resonator. The waveguide could be used to read out the rotational state of the ring.<sup>707</sup> Finally, as shown in Fig. 59(d), one could use a TAAP ring to couple two dipole waveguides to each other in a velocity selective fashion, much like wavelength selective multiplexing using tunable whispering gallery resonators.<sup>708,709</sup>

## B. Ring-leads system

A widely studied system within mesoscopic physics are rings attached to leads with an applied magnetic field.<sup>698,710</sup> This system features the Aharonov–Bohm effect, where currents through the ring can be modulated with an applied magnetic field. While extensively studied for fermions such as electrons, it is not well understood for bosonic type of systems. Atomtronic setups allow for the study of these types of systems in a controlled way for the first time. A theoretical study has been performed in Refs. 690 and 690, which is reviewed in this section. Atomtronic setups for transport can be modeled using the Bose–Hubbard model.<sup>125</sup> Here, circuits of ring-leads or Y-junctions as seen in Fig. 60 are modeled as a lattice system. For example, for the ring-leads system, the three individual components (source lead, ring, and drain) are each modeled as one-dimensional chains with nearest-neighbor tunneling interactions. The different components are then coupled together via tunneling at specific lattice sites. A ring with an even number of lattice sites  $L$  coupled to two leads [see Fig. 60(b)] is given by the Hamiltonian  $\mathcal{H} = \mathcal{H}_r + \mathcal{H}_l$ . The ring part of the Hamiltonian is given by

$$\mathcal{H}_r = - \sum_{j=0}^{L-1} \left( J e^{i2\pi\Phi/L} \hat{a}_j^\dagger \hat{a}_{j+1} + \text{h.c.} \right) + \frac{U}{2} \sum_{j=0}^{L-1} \hat{n}_j (\hat{n}_j - 1), \quad (105)$$

where  $\hat{a}_j$  and  $\hat{a}_j^\dagger$  are the annihilation and creation operator at site  $j$ ,  $\hat{n}_j = \hat{a}_j^\dagger \hat{a}_j$  is the particle number operator,  $J$  is the intra-ring hopping,  $U$  is the on-site interaction between particles, and  $\Phi$  is the total flux through the ring. Periodic boundary conditions are applied:  $\hat{a}_L^\dagger = \hat{a}_0^\dagger$ . The two leads dubbed source (S) and drain (D) consist of a single site each, which are coupled symmetrically at opposite sites to the ring with coupling strength  $K$ . In both of them, local potential energy and on-site interaction are set to zero as the leads are considered to be large with low atom density. The lead Hamiltonian is  $\mathcal{H}_l = -K(\hat{a}_S^\dagger \hat{a}_0 + \hat{a}_D^\dagger \hat{a}_{L/2} + \text{h.c.})$ , where  $\hat{a}_S^\dagger$  and  $\hat{a}_D^\dagger$  are the creation operators of source and drain, respectively.

The dynamics of transport within this system in various settings is discussed in Refs. 689 and 690, which we now review. A way to probe transport was studied in the case where the source and drain consist of only one lattice site each. Here, the atoms are initially prepared in the source with the ring and drain being empty of atoms. During the time evolution, the atoms flow out of the source lead and propagate via the ring to the drain. In the weak-coupling regime  $K/J \ll 1$ , the lead-ring tunneling is slow compared to the dynamics inside the ring [see Figs. 59(a), 59(c), and 59(e)]. In this regime, the condensate mostly populates the drain and source, leaving the ring nearly empty. As a result, the scattering due to the on-site interaction  $U$  has a negligible influence on the dynamics. With increasing flux  $\Phi$ , the oscillation becomes faster and the ring populates, resulting in

increased scattering and washed-out density oscillations. In the strong-coupling regime  $K/J \approx 1$ , the lead-ring and the intraring dynamics are characterized by the same frequency and cannot be treated separately. Here, a superposition of many oscillation frequencies appears, and after a short time, the condensate is evenly spread in both leads and ring [Figs. 59(b), 59(d), and 59(f)]. The density in the ring is large and scattering affects the dynamics by washing out the oscillations. Close to  $\Phi = 0.5$ , the oscillations slow down, especially for weak interaction, due to destructive interference.<sup>711</sup>

The authors also studied the dynamics for the case where the source and drain leads consists of many sites, probing the regime of many atoms in an extended system. Here, the large source lead is filled with atoms and then suddenly coupled to the ring to generate the dynamics. The authors model this in two ways: In the first case, they solve the full dynamics of leads and ring using DMRG.<sup>712,713</sup> Then, they study an approximate method, where the leads are approximated as a large bath and are traced out. The resulting dynamics is described using the Lindblad master equation,

$$\frac{\partial \rho}{\partial t} = -\frac{i}{\hbar} [H, \rho] - \frac{1}{2} \sum_m \left\{ \hat{L}_m^\dagger \hat{L}_m, \rho \right\} + \sum_m \hat{L}_m \rho \hat{L}_m^\dagger,$$

for the reduced density matrix<sup>714</sup> within the Born–Markov approximation with  $L_1 = \sqrt{\Gamma} \hat{a}_S^\dagger$ ,  $L_2 = \sqrt{\Gamma} \hat{a}_S$ , and  $L_3 = \sqrt{\Gamma} \hat{a}_D$  ( $r$  characterizes the back-tunneling into the source reservoir). Then, the steady state of the density matrix is solved  $\partial \rho_{SS} / \partial t = 0$ .<sup>715</sup> The operator for the current is  $j = -iK(\hat{a}_S^\dagger \hat{a}_0 - \hat{a}_0^\dagger \hat{a}_S)$ , and its expectation value is  $\langle j \rangle = \text{Tr}(j \rho_{SS})$ . In Figs. 62(a)–62(c), the authors compare the open system Lindblad approach with a full simulation of both ring and reservoirs using DMRG.<sup>712,713</sup> Both methods yield similar results with the Lindblad approximation smoothing out the oscillation found in DMRG. This shows that leads modeled as Markovian bath without memory are sufficient to describe the dynamics. Further, they investigate the evolution of the current toward the steady-state. They find that the initial dynamics depends on the flux, which is a sign of the Aharonov–Bohm effect being initially present. However, the steady-state reached after long times is nearly independent of flux, demonstrating the absence of the Aharonov–Bohm effect in the steady state for interacting bosons.

## C. Y-junctions

The Y-junction is a system consisting of three one-dimensional chains, which are coupled together at a single point [see Fig. 60(c)]. They have been of wide interest in mesoscopic physics for electronic type systems.<sup>716</sup> For cold atoms, such systems have been proposed and realized experimentally.<sup>28,152,155,717–719</sup> In Ref. 690, the authors studied theoretically the dynamics of density wave excitations that are transmitted and reflected in a cold atom Y-junction. They are interested in how the system evolves for varying the coupling strength of the Y-junction. They find characteristic regimes of transmission and reflections.

The Hamiltonian for the Y-junction is  $\mathcal{H}_S + \mathcal{H}_D + \mathcal{H}_I$  with the source lead Hamiltonian (analog for the two drain leads),

$$\mathcal{H}_S = - \sum_{j=1}^{L_S-1} \left( J \hat{s}_j^\dagger \hat{s}_{j+1} + \text{h.c.} \right) + \sum_{j=1}^{L_S} \frac{U}{2} \hat{n}_j^s (\hat{n}_j^s - 1), \quad (106)$$

where  $\hat{s}_j$  and  $\hat{s}_j^\dagger$  are the annihilation and creation operators at site  $j$  in the source lead,  $\hat{n}_j^s = \hat{s}_j^\dagger \hat{s}_j$  is the particle number operator of the source,  $J$  is the intra-lead hopping,  $L_S$  is the number of source lead sites, and  $U$  is the on-site interaction between particles. All units are rescaled in terms of the hopping term  $J$ . The Hamiltonian  $\mathcal{H}_D$  for the two drain leads have similar Hamiltonians, where one replaces the index  $s$  with respective  $d$  (for first drain) and  $f$  (second drain) and defines the drain length  $L_D$ . The coupling Hamiltonian between the source lead and the two drain leads is

$$\mathcal{H}_I = -K\hat{s}_1^\dagger(\hat{d}_1 + \hat{f}_1) + \text{h.c.}, \quad (107)$$

where  $K$  is the coupling strength between source and drain leads. The current through the Y-junction is defined as

$$j_Y = -iK\hat{s}_0^\dagger\hat{d}_0 + \text{h.c.} \quad (108)$$

To study the propagation of a density excitation through the setup, the authors prepare the system in the ground state of the full Hamiltonian with initially a small local potential offset in the lead Hamiltonian. This will create a localized density bump in the source lead. Then, they add the following Hamiltonian for the offset potential to the source Hamiltonian,

$$\mathcal{H}_P = -\varepsilon_D \sum_{j=1}^{L_S} \exp\left(-\frac{(j-d)^2}{2\sigma^2}\right) \hat{n}_j^s, \quad (109)$$

where  $d$  is the distance of the initial excitation to the junction. At the start of the time evolution, the offset potential is instantaneously switched off. The density bump will propagate as a density excitation in both positive and negative direction, where here only the forward direction is regarded.

The authors develop a method to calculate the amount of transmitted and reflected density waves. They calculate the total density of the incoming wave by taking the first  $a$  sites of the source lead at a specific time  $t_{\text{in}}$  when the density waves has entered this region, and subtracting from it the density at time  $t=0$  before the wave has entered the region,

$$N_{\text{inc}} = \sum_{i \in a \text{ sites of source}} [n_i(t_{\text{in}}) - n_i(0)]. \quad (110)$$

Here,  $n_i(t)$  is the expectation value of the density at the  $i$ -th site of the system at time  $t$ . The transmission coefficient is found by dividing the change in the atom number in the drain density by the total density of the incoming wave

$$T = \frac{\sum_{i \in \text{drain}} [n_i(t) - n_i(0)]}{N_{\text{inc}}}, \quad (111)$$

and the reflection coefficient as

$$R = 1 - T. \quad (112)$$

The limiting cases of the infinitely strong on-site interaction with hard-core bosons are presented in Fig. 64. In Figs. 64(a) and 64(c), the authors study the propagation for different values of lead coupling  $K$ .

In the source lead, an initial excitation is prepared as a small patch of increased density. At  $t=0$ , the potential offset is quenched, and the excitation starts moving in the forward and backward directions. The forward moving part of the wave propagates from the source through the junction to the two drain leads. At the junction between the chains (site 160 in the graph), the wave is both transmitted and reflected. For the reflection amplitude, three characteristic reflection regimes are found, which are controlled by the junction coupling  $K$ .

First, look at the reflection peak as seen in Fig. 64(e) at time  $tJ = 27$ . In the strong coupling regime  $K = 1J$ , a negative (Andreev-like) reflection amplitude peak is found. For the intermediate coupling regime  $K \approx 0.5$ , the back reflection amplitude is very small, and the reflected wave consists of a small, first positive and then negative part, of nearly equal weight. Finally, for the weak coupling regime with  $K$  small, a large positive back-reflection with small transmission is found. In Table II, the total transmitted and reflected density at time  $t = 31/J$  is calculated using Eqs. (110)–(112). This gives the transmission and reflection coefficient of the density wave packet. For strong coupling  $K = 1J$  with Andreev reflections, the transmission coefficient is nearly  $T \approx 4/3$ , which corresponds to the theoretical value predicted for a Y-junction in the limit of weak interaction, within the Gross-Pitaevskii equation.<sup>715</sup> In this regime, the transmission is larger than the initial density wave, owing to the negative reflection  $R \approx -1/3$ . Similar dynamics is also found also for finite interaction  $U$  within the Bose-Hubbard model.<sup>690</sup>

#### D. Differences between fermions and hard-core boson

Bosons and fermions differ fundamentally in their particle exchange relations: The bosonic many-body wavefunction is symmetric, while fermions are antisymmetric under exchange of two particles. As a result of these properties, the Pauli principle is enforced for fermions: at a single site, only zero or one fermion can exist, while noninteracting bosons do not have this restriction. However, for strongly interacting bosons in the hard-core limit, only one hard-core boson can be at a single site, mimicking the Pauli principle, while maintaining a symmetric many-body wavefunction. In one dimension, hard-core bosons and fermions are equivalent and a mapping between fermions and hard-core bosons exists; however, this is not the case beyond one-dimensional systems. The effect of this feature on transport has been studied numerically in detail in Ref. 690.

In a Y-junction with the same setup as in Sec. XII C, fermions and hard-core bosons show fundamentally different types of reflection behavior. Figure 65 shows the density wave for transmission and reflection for both types of particles. Hard-core bosons show a clear Andreev-reflection with negative reflection, while spinless fermions do not.

Similar differences arise in the ring-lead system. In a half-filled system, a density wave is excited similar to procedure detailed earlier

TABLE II. Transmission and reflection coefficients calculated at  $t = 31/J$  with Eqs. (110)–(112) ( $t_{\text{in}} = 15$ ,  $a = 30$ ).

|              | $K = 1$ | $K = 0.5$ | $K = 0.2$ |
|--------------|---------|-----------|-----------|
| Transmission | 1.332   | 0.947     | 0.207     |
| reflection   | -0.332  | 0.053     | 0.793     |



introduced in Sec. XII C. The reflected and transmitted density wave for zero and half-flux is shown in Fig. 63. First, the authors study the properties of the reflected density wave. For zero flux, the reflected density wave is different for fermions and bosons. For hard-core bosons, the same characteristic Andreev-like negative reflection peak as seen in the strongly coupled Y-junction. However, the reflection dynamics for hard-core bosons is flux independent. Now, observe the transmission (upper curve). Spinless fermions become flux dependent. Here, the fermionic density waves are transmitted for zero flux, while at half-flux, zero transmission is observed due to Aharonov–Bohm interference. However, for hard-core bosons, the transmission is unaffected by flux. This is again demonstrating the absence of the Aharonov–Bohm effect for bosons. In short, density excitations for fermions show the Aharonov–Bohm effect, while for interacting bosons, the Aharonov–Bohm effect is absent.

### E. Entangled state generation with topological pumping in ring circuits

Directed transport can be also engineered by periodically driving a system. Here, topological pumping, pioneered by Thouless,<sup>700,701</sup> can transport excitations with the added feature that the transport is protected against noise and imperfections due to the topological properties of the system. This has been successfully demonstrated with cold atoms.<sup>702,720</sup> Topological pumping is realized adiabatically by driving the parameters of Hamiltonians with topological features periodically in time. This idea can be extended to interacting many-body systems.<sup>703,704</sup> This type of driving can be extended to transport atoms through ring-lead circuits as shown in Ref. 691 with a similar setup as introduced for the ring-lead circuit (see Sec. XII B). To enable pumping, a time-dependent and spatially varying local potential is added to the system, which is modulated periodically. If engineered correctly, it pumps atoms from the source lead, through the ring, into the drain. For the exact details on the procedure, refer to Ref. 691.

Here, we will shortly review a key result of this study: Applying topological pumping to ring-lead systems can be used to create highly entangled quantum states.  $N$  atoms are initially placed in the source lead. The pumping is switched on, and the pumping transfers particles into the ring. At the junction of source lead and ring, two possible directions along the ring open up: Either going along the top or bottom path of the ring. Here, the Bose–Hubbard interaction term  $U$  can lead to the creation of NOON-like superposition states, where  $N$  atoms either go along either of the two paths  $[(\Psi_{\text{NOON}}) = \frac{1}{\sqrt{2}}(|0\rangle \otimes (|N0\rangle + |0N\rangle)]$ . This concept can be exemplified by a simplified three site system, consisting of a single source lead site, which is coupled two other sites which represents a part of the ring. The fidelity  $F = |\langle \Psi_{\text{NOON}} | \Psi \rangle|^2$  of the creation of the NOON like entangled state is plotted in Fig. 66. For a particular set of parameters, NOON states of up to six particles with nearly unit fidelity can be created. For more particles or higher interaction, the fidelity decreases due to an exponential suppression of the energy gap. This setup is a powerful method to prepare and study highly entangled states of cold atoms.

### F. Concluding remarks and outlook

Transport in quantum many-body systems is a fundamental problem important for quantum information and condensed matter physics. Cold atoms can be used to simulate these transport problems

in novel regimes that are difficult to realize within other setup. With recent technologies, transport with nearly no heating can be achieved by extremely smooth atom waveguides. Further, new regimes of transport can be studied by attaching reservoirs of atoms to the system that induce a current through the system. Studying the current can reveal properties of the system that are hard to extract otherwise.

TAAP rings and engineered waveguides made of light allow construction of atomic circuits that can transport cold atoms in various configurations. Here, we reviewed the properties of two particular atomic circuits that are enabled by this technology, namely, ring-lead systems and Y-junctions.<sup>689,690</sup> The current through these circuits holds some surprises: Bosonic Y-junctions show Andreev-reflections, known from fermionic superconductor-metal interfaces. By tuning the coupling of the Y-junction, the type of reflection can be tuned between regular (positive) and negative Andreev reflections. These Andreev reflection are well known from electronic-superconducting interfaces, as such it is surprising to observe them in a bosonic system as well. For transport through ring-lead systems, the current for interacting bosons is independent of flux and the Aharonov–Bohm effect is absent. This is in stark contrast to fermionic systems, which are highly flux dependent as shown in electronic systems. This difference allows one to study the fundamental difference between fermions and bosons due to their antisymmetric and symmetric many-body wavefunction in a transport experiment, simply by studying the current of the system. Finally, by changing the circuit potential in time, topological pumping can be realized to transport atoms in a robust fashion and create highly entangled states of NOON-type. These states could become very useful for quantum-enhanced sensing as the sensitivity of NOON-states increases linearly with the number of entangled particles. For example, the NOON states could be applied to sense rotation. While the atoms are pumped through the ring, they pick up a phase that is proportional to the rotation affecting the ring times the number of atoms.

In a very interesting future direction transport through nonstandard type Hubbard models could be investigated. These types of Hubbard models feature higher order tunneling and interactions terms that create novel effects and phases. These types of Hamiltonians can nowadays be realized within cold atoms experiments.<sup>721</sup>

The proposed setups can be realized in state-of-the art experiments with both bosonic and fermionic cold atoms. The confinement for the atoms in the form of the circuits can be constructed using DMDs or other light-based potential painting techniques, allowing for arbitrary potential shapes and even time-dependent modulation of the potential. While engineering more complex potentials and driving protocols may be a time-consuming task for humans, new machine learning methods could help us to engineer improved potentials and cold atom dynamics automatically without human intervention.<sup>722</sup>

## XIII. ARTIFICIAL QUANTUM MATTER IN LADDER GEOMETRIES

**V. Ahufinger, R. Citro, S. De Palo, A. Minguzzi, J. Mompert, E. Orignac, N. Victorin**

The fractional quantum Hall effect<sup>723,724</sup> is a striking example of the interplay of interaction and topology in condensed matter physics. It is characterized by many fascinating properties such as a precise quantization of the Hall resistance depending only on fundamental constants, excitations carrying fractional charges with anyonic

statistics, and dissipationless chiral edge modes. While the effect has been initially observed with fermions, bosonic analogs have been proposed by Regnault and Jolicoeur in rotating clouds of ultracold atoms.<sup>725</sup> Recently, the realization in experiments of artificial gauge fields<sup>591,726–728</sup> has opened another route for observing quantum Hall phases with ultracold atoms. As a first step toward the realization of quantum Hall phases with ultracold atoms, it is interesting to consider the so called ladder systems,<sup>729–753</sup> i.e., two dimensional systems that are of finite size along one of the dimensions. Such a deceptively simple system is already sensitive to the effect of the applied flux and can exhibit analogs of the quantum Hall phase.<sup>737,738,747</sup> Moreover, it shows a wealth of phases, emerging from interplay of rung and leg tunnel, interactions, artificial gauge field, filling.<sup>732,733,737,751,752</sup> For bosonic atoms, in low flux, an analog of the Meissner phase is obtained.<sup>729,730</sup> At high flux, a quasi-long range ordered vortex phase is formed.<sup>729,730</sup> Interleg interactions can stabilize an atomic density wave at intermediate flux.<sup>752,753</sup> For a flux commensurate with the density, the analog of QHE is found.<sup>737,738,747</sup> At a different commensuration between flux and density, an incommensuration driven by interchain hopping is obtained.<sup>750</sup> Furthermore, a variant of the ladder in the form of a diamond chain has topological properties<sup>419</sup> and allows one to simulate quantum magnetism.<sup>754,755</sup>

## A. A boson ring ladder at weak interactions

### 1. Model

We consider  $N$  bosons occupying two coupled one-dimensional concentric lattice rings subjected to two artificial gauge fields ad organized on a planar geometry. The stacked geometry has also been thoroughly studied.<sup>580,692</sup> This system could be experimentally realized, e.g., using dressed potentials,<sup>111</sup> or Laguerre–Gauss beams.<sup>140</sup> The Hamiltonian reads

$$\begin{aligned}\hat{H} &= \hat{H}_0 + \hat{H}_{int} \\ &= -J \sum_{l=1, p=1,2}^{N_i} \left( b_{l,p}^\dagger b_{l+1,p} e^{i\Phi_p} + b_{l+1,p}^\dagger b_{l,p} e^{-i\Phi_p} \right) \\ &\quad - \frac{\Omega}{2} \sum_{l=1}^{N_i} \left( b_{l,1}^\dagger b_{l,2} + b_{l,2}^\dagger b_{l,1} \right) + \frac{U}{2} \sum_{l=1, p=1,2}^{N_i} b_{l,p}^\dagger b_{l,p}^\dagger b_{l,p} b_{l,p}, \quad (113)\end{aligned}$$

where  $b_{l,p}$  are the bosonic field operators for the  $p$ -th ring,  $l$  indicates the site position on each ring made of  $N_i$  sites,  $J$  is the tunneling amplitude along each ring, threaded by the fluxes  $\Phi_{1,2}$ , respectively, and  $\Omega$  is the inter-ring tunneling amplitude. In the noninteracting regime  $U = 0$ , this model is readily diagonalized, yielding a two-band excitation spectrum

$$\begin{aligned}E_{\pm}(k) &= -2J \cos(\lambda/2) \cos(k - \Phi) \\ &\quad \pm \sqrt{(\Omega/2)^2 + (2J)^2 \sin(\lambda/2)^2 \sin(k - \Phi)^2}, \quad (114)\end{aligned}$$

where we have set  $\lambda = \Phi_1 - \Phi_2$  and average flux  $\Phi = (\Phi_1 + \Phi_2)/2$ . Depending on the ratio  $\Omega/J$  and on  $\lambda$ , the lowest band of the excitation spectrum has either one or two minima centered at  $k = \Phi$ . The ground state of the Bose gas is a Bose–Einstein condensate occupying the minima of such an excitation spectrum. In the case of a single minimum, the ground state corresponds to the Meissner phase, and in the

case of a coherent superposition of the occupancy of the two minima, the ground state is in the vortex phase. The Meissner phase is characterized by vanishing transverse current and homogeneous density profile. The vortex phase has nonzero transverse current and density modulations along the ring. The vortex to Meissner phase transition has been experimentally observed in Ref. 756. The chiral current on the ring, i.e., the difference of longitudinal currents among the two rings, is characterized by subsequent jumps each time a vortex enters into the system.

### 2. Ground state of weakly interacting ring

We assume large occupancy of the lattice sites and weak interactions  $U/J \ll 1$ . In this regime, we describe the system by the mean-field approximation. Setting  $\Psi_{l,p}(t) = \langle b_{l,p}(t) \rangle$ , the condensate wavefunction, we solve the coupled discrete nonlinear Schroedinger equations (DNLS)

$$\begin{aligned}i\partial_t \Psi_{l,1}(t) &= -J \Psi_{l+1,1}(t) e^{i(\Phi+\lambda/2)} - J \Psi_{l-1,1}(t) e^{-i(\Phi+\lambda/2)} \\ &\quad - K \Psi_{l,2}(t) + U |\Psi_{l,1}(t)|^2 \Psi_{l,1}(t), \quad (115)\end{aligned}$$

$$\begin{aligned}i\partial_t \Psi_{l,2}(t) &= -J \Psi_{l+1,2}(t) e^{i(\Phi-\lambda/2)} - J \Psi_{l-1,2}(t) e^{-i(\Phi-\lambda/2)} \\ &\quad - K \Psi_{l,1}(t) + U |\Psi_{l,2}(t)|^2 \Psi_{l,2}(t). \quad (116)\end{aligned}$$

We have also assumed that the interaction energy is smaller than the bandgap such that the single-band approximation for each ring lattice holds. At varying  $UN/JN_i$  and  $\Omega/J$ , the ground state displays three phases:<sup>419</sup> the vortex (V) and Meissner (M) phases found in the noninteracting regime as well as the biased-ladder phase characterized by imbalanced density populations among the two rings and uniform density profile. We denote the latter (BL-V) or (BL-M) as shown in Fig. 67, depending whether for the same values of  $\lambda$  and  $\Omega/J$ , the corresponding noninteracting spectrum has one or two minima. Notice that the BL phases are only found at weak interactions, while they are disrupted as the interactions increase.<sup>757</sup> For this reason, they are not found in the DMRG calculations of Sec. XIII B.

### 3. Excitation spectrum of weakly interacting ring

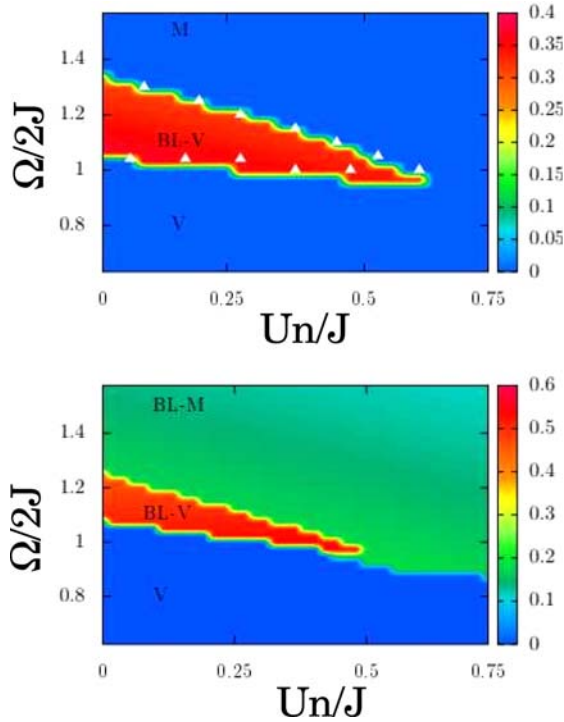
We next present our results for the excitation spectrum of the weakly interacting Bose gas on a ring lattice.<sup>758</sup> Within the Bogoliubov approximation, we set  $\hat{a}_{l,p} = \Psi_{l,p}^{(0)} + \delta \hat{a}_{l,p}$ , where  $\Psi_{l,p}^{(0)}$  is the ground state solution with chemical potential  $\mu$ . we find the excitation spectrum using the expansion of the fluctuation operator  $\delta \hat{a}_{l,p}$  in normal modes with energy  $\omega_\nu$ , according to

$$\delta \hat{a}_{l,p} = \sum_{\nu} h_{\nu,l}^{(p)} \hat{\gamma}_{\nu} - Q_{\nu,l}^{*(p)} \hat{\gamma}_{\nu}^{\dagger}. \quad (117)$$

The solution Bogoliubov–de Gennes eigenvalue equations for the mode amplitudes  $h_{\nu,l}^{(p)}$  and  $Q_{\nu,l}^{(p)}$  yields the excitation spectrum. We use both eigenvalues and eigenvectors of the Bogoliubov equations to compute the dynamic structure factor

$$S_{p,p'}(q, \omega) = \sum_{s \neq 0} |\langle s | \hat{\rho}_q^{(p,p')} | 0 \rangle|^2 \delta(\omega - \omega_s). \quad (118)$$

As an example, we show the results for the excitation spectrum in the Meissner and in the vortex phase. The excitation spectrum is strongly dependent on the phase of the underlying ground state. In the



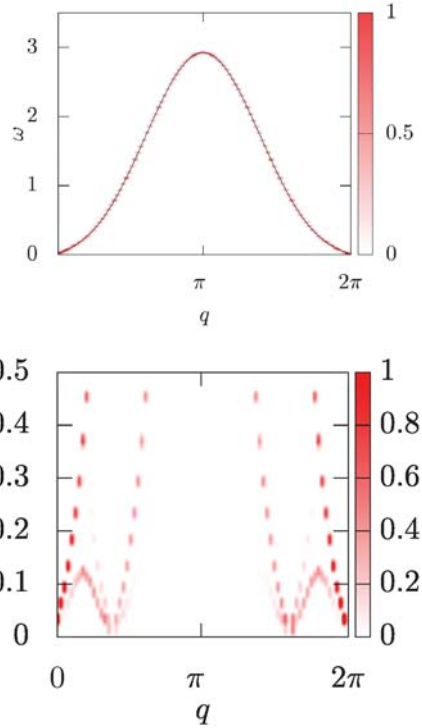
**FIG. 67.** Color map of the imbalance among particle numbers in each ring, in the  $(\Omega/J, UN/JN_s)$  plane, for (upper panel)  $\lambda = \pi/2$ ,  $\Phi = 6\pi/N_s$  and  $N_s = 20$ , (lower panel)  $\lambda = \pi/2$ ,  $\Phi = \pi/N_s$  and  $N_s = 20$ . The letters indicate the parameter regimes where we find a biased-ladder phase (BL-V) where the single-particle spectrum has a double minimum, a Meissner phase (M), a vortex phase (V) and a biased-ladder phase (BL-M) where the single-particle spectrum has a single minimum. White triangles represent the frontiers between biased-ladder phase the two other phase, namely, vortex phase and Meissner phase as calculated with the variational ansatz including finite size effect.

Meissner phase, a single Goldstone mode is found and corresponds to the  $U(1)$  symmetry breaking associated with the formation of a Bose–Einstein condensate. In the vortex phase, two Goldstone modes are observed. Indeed in the vortex phase, a second symmetry is broken, i.e., the discrete translational symmetry. We have shown that this is associated with the emergence of supersolidity of the gas and is corroborated by the calculation of the static structure factor, displaying a well-defined peak as shown in Fig. 68, and the first-order correlation function, demonstrating phase coherence.

### B. The boson ladder at strong interaction

Let us consider a spin-1/2 bosons with spin-orbit interaction model,<sup>751</sup> where  $\Omega$  is the transverse magnetic field,  $\lambda$  the spin-orbit coupling,  $U_{\uparrow\uparrow} = U_{\downarrow\downarrow} = U$  the repulsion between bosons of identical spins,  $U_{\uparrow\downarrow} = U_{\perp}$  the interaction between bosons of opposite spins. Its Hamiltonian is<sup>745,747</sup>

$$H = -J \sum_{j,\sigma} (b_{j,\sigma}^\dagger e^{i\lambda\sigma} b_{j+1,\sigma} + b_{j+1,\sigma}^\dagger e^{-i\lambda\sigma} b_{j,\sigma}) + \frac{\Omega}{2} \sum_{j,\alpha,\beta} b_{j,\alpha}^\dagger (\sigma^x)_{\alpha\beta} b_{j,\beta} + \sum_{j,\alpha,\beta} U_{\alpha\beta} n_{j,\alpha} n_{j,\beta}, \quad (119)$$



**FIG. 68.** Dynamic structure factor in the frequency-wavevector plane in the frequency-wavevector plane (color map,  $q$  in units of  $1/a$  with a lattice spacing and  $\omega$  in units of  $J$ ) Upper panel: in the Meissner phase, for  $Un/J = 0.2$ ,  $\lambda = \pi/2$ ,  $K/J = 3$ . Lower panel: in the vortex phase for  $\Omega/J = 1.6$ ,  $\lambda = \pi/2$ ,  $Un/J = 0.2$ . For both panels  $N_s = 80$ .

where  $\sigma = \pm 1/2$  or  $\sigma = \uparrow, \downarrow$  is the spin index<sup>677,759,760</sup> and  $j$  is the site index,  $n_{j\alpha} = b_{j\alpha}^\dagger b_{j\alpha}$ . The Hamiltonian [Eq. (119)] is mapped onto the Hamiltonian of the two-leg ladder in flux [Eq. (113)] by making  $p = \frac{3}{2} - \sigma$ ,  $\Phi_p = (-)^{p-1} \lambda/2$ ,  $U_{\perp} = 0$ .

The low-energy effective theory for the Hamiltonian [Eq. (119)], treating  $\Omega$  and  $U_{\perp}$  as perturbations, can be obtained by using Haldane's bosonization of interacting bosons.<sup>761</sup> Introducing<sup>761</sup> the fields  $\phi_\alpha(x)$  and  $\Pi_\alpha(x)$  satisfying canonical commutation relations  $[\phi_\alpha(x), \Pi_\beta(y)] = i\delta(x-y)$  as well as the dual  $\theta_\alpha(x) = \pi \int^x dy \Pi_\alpha(y)$  of  $\phi_\alpha(x)$ , and after introducing the respective combinations of operators  $\phi_{c,s} = (\phi_{\uparrow} \pm \phi_{\downarrow})/\sqrt{2}$ , we can represent the low-energy Hamiltonian as  $H = H_c + H_s$ , where

$$H_c = \int \frac{dx}{2\pi} \left[ u_c K_c (\pi \Pi_c)^2 + \frac{u_c}{K_c} (\partial_x \phi_c)^2 \right] \quad (120)$$

describes the total density fluctuations for incommensurate filling when umklapp terms are irrelevant, and

$$H_s = \int \frac{dx}{2\pi} \left[ u_s K_s \left( \pi \Pi_s + \frac{\lambda}{a\sqrt{2}} \right)^2 + \frac{u_s}{K_s} (\partial_x \phi_s)^2 \right] - 2\Omega a_0^2 \int dx \cos \sqrt{2}\theta_s + \frac{U_{\perp} a B_1^2}{2} \int dx \cos \sqrt{8}\phi_s \quad (121)$$



describes the antisymmetric density fluctuations. In Eqs. (121) and (120),  $u_s$  and  $u_c$  are, respectively, the velocity of antisymmetric and total density excitations, and  $A_0$  and  $B_1$  are nonuniversal coefficients,<sup>762</sup> while  $K_s$  and  $K_c$  are the corresponding Tomonaga–Luttinger (TL) exponents.<sup>753</sup> They can be expressed as a function of the velocity of excitations  $u$  and Tomonaga–Luttinger liquid exponent  $K$  of the isolated chain.<sup>753</sup>

For an isolated chain of hard core bosons, we have  $u = 2J \sin(\pi\rho_0)$  and  $K = 1$ . The phase diagram of the Hamiltonian can be determined by looking at physical observables such as the rung and leg current, momentum distribution, and correlation functions. Physical observables can be all represented in bosonization. The rung current, or the flow of bosons from the upper leg to the lower leg, is

$$J_{\perp}(j) = -i\Omega(b_{j,\uparrow}^{\dagger}b_{j,\downarrow} - b_{j,\downarrow}^{\dagger}b_{j,\uparrow}) \\ = 2\Omega A_0^2 \sin \sqrt{2}\theta_s + \dots \quad (122)$$

The chiral current, i.e., the difference between the currents of upper and lower leg, is defined as

$$J_{\parallel}(j, \lambda) = -ij \sum_{\sigma} \sigma (b_{j,\sigma}^{\dagger} e^{i\lambda\sigma} b_{j+1,\sigma} - b_{j+1,\sigma}^{\dagger} e^{-i\lambda\sigma} b_{j,\sigma}) \quad (123)$$

$$= \frac{u_s K_s}{\pi\sqrt{2}} \left( \partial_x \theta_s + \frac{\lambda}{a\sqrt{2}} \right). \quad (124)$$

The density difference between the chains  $S_j^z = n_{j\uparrow} - n_{j\downarrow}$  is written in bosonization as

$$S_j^z = -\frac{\sqrt{2}}{\pi} \partial_x \phi_s - 2B_1 \sin \left( \sqrt{2}\phi_c - \pi\rho x \right) \sin \sqrt{2}\phi_s, \quad (125)$$

while the density of particles per rung is

$$n_j = -\frac{\sqrt{2}}{\pi} \partial_x \phi_c - 2B_1 \cos \left( \sqrt{2}\phi_c - \pi\rho x \right) \cos \sqrt{2}\phi_s. \quad (126)$$

When  $\Omega \neq 0$ ,  $U_{\perp} = 0$ , and  $\lambda \rightarrow 0$ , the antisymmetric modes Hamiltonian Eq. (121) reduces to a quantum sine-Gordon Hamiltonian. For  $K_s > 1/4$ , the spectrum of  $H_s$  is gapped and the system is in the so-called Meissner state<sup>729,730</sup> characterized by  $\langle \theta_s \rangle = 0$ . In such a state, the chiral current increases linearly with the applied flux at small  $\lambda$ , while the average rung current  $\langle J_{\perp} \rangle = 0$  and its correlations  $\langle J_{\perp}(j)J_{\perp}(0) \rangle$  decay exponentially with distance. The transition from the Meissner to the Vortex phase is signaled by the splitting of the momentum distribution  $n(k)$  from  $k=0$  to a finite  $Q = \sqrt{\lambda^2 - \lambda_c^2}$  that depends on the spin–orbit interaction  $\lambda$ . For this reason, the transition falls into the universality class of the commensurate–incommensurate transitions (C–IC). The rung current correlation function develops two symmetric peaks, and the spin static structure is linear at low momentum. The phase diagram for a hard-core bosonic ladder at  $n=1$ , obtained using density renormalization group (DMRG) technique,<sup>763,764</sup> is shown in Fig. 69 together with the momentum distribution  $n(k)$  as a function of  $\lambda$  across the C–IC transition. Compared to the noninteracting case, the phase diagram as a function of  $\Omega/J$  and  $\lambda$  shows an enlargement of the Meissner phase and its persistence above a certain value of  $\Omega/J$ . Above a certain value

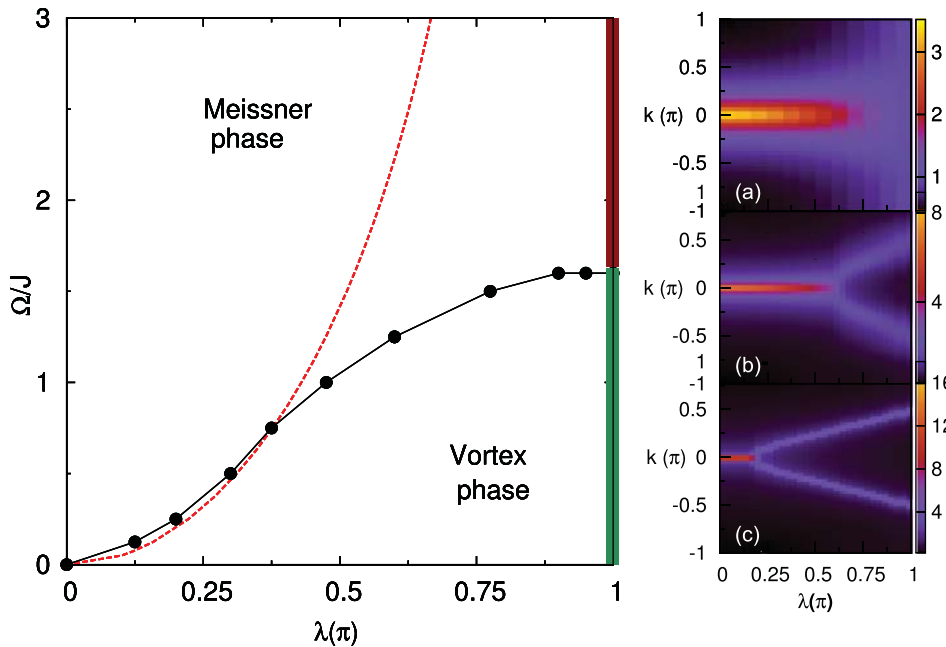
of  $\lambda$ , a second incommensuration appears in the rung current correlation functions and the static structure factor. Such incommensuration is associated with the appearance of an extra peak in the rung current correlation function at wavevectors  $P = \sqrt{\lambda^2 + p(\Omega^2)}$  and  $\pi \pm P$  with  $p(\Omega)$  a function of the interchain tunneling. For  $\lambda = \pi$ , the correlation functions show a tendency to a localized regime.

With  $\Omega \neq 0$ ,  $U_{\perp} \neq 0$ , the C–IC transition is replaced by a Meissner-to-incommensurate charge density wave (ICDW) which falls into the Ising universality class, followed by a melting of the Vortex phase at large enough  $\lambda$ , going toward a BKT transition when entering the Vortex phase.<sup>753</sup> The melting of the Vortex phase is signaled by the Lorentzian shape of the momentum distribution peaks preceded by a Lifshitz point.<sup>765</sup> In the phase diagram, see Fig. 70 for the case of a hard-core bosonic ladder in the presence of an attractive interaction  $U_{\perp}$ , obtained using DMRG simulations, it is possible to trace these features. As we increase the interaction strength [panel (a) in Fig. 70], the charge structure factor develops peaks at  $k = \pi/2$  and  $k = 3\pi/2$  and  $S_s(k)$  has an almost quadratic behavior at small wave vectors. The quadratic behavior indicates that spin excitations remain gapped, while the presence of peaks at  $k = \pi/2, 3\pi/2$  in  $S_c(k)$  is the signature of a zigzag charge density wave. Going to panel (b), at increasing  $\lambda$ , the momentum distribution develops two broad maxima indicating a vortex melted phase, while the rung current correlation function starts to form two bumps. Increasing still  $\lambda$ , in panel (c) both the momentum distribution, as well the rung-current correlation function  $C(k)$ , develop two separate peaks,<sup>730</sup> that show negligible size effects, indicating the presence of an incommensuration. Conversely both the charge and spin structure factors have a linear behavior at small  $k$  indicating a gapless phase.

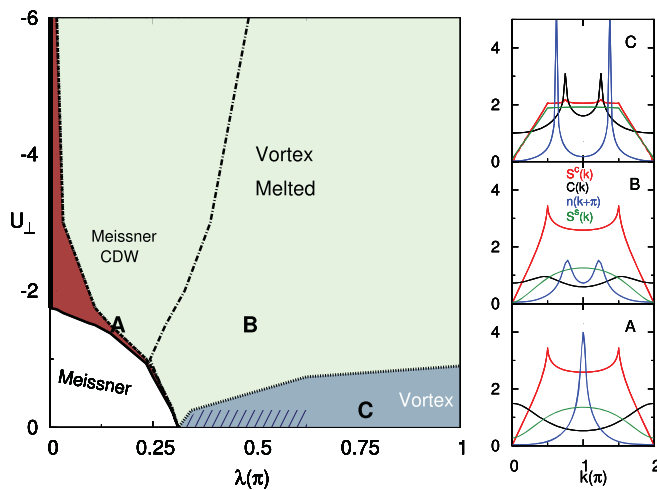
## C. Ultracold atoms carrying orbital angular momentum (OAM) in a diamond chain

### 1. Topological edge states and Aharonov–Bohm caging

We consider a ladder with a diamond-chain shape with a unit cell formed by three cylindrically symmetric potentials of radial frequency  $\omega$ , forming a triangle with central angle  $\Theta$  and nearest-neighbor separation  $d$  (see Fig. 71). Noninteracting ultracold atoms of mass  $m$  that may occupy the two degenerate OAM  $l=1$  states with positive or negative circulation localized at each site are loaded into the ladder. Such a system could be experimentally implemented, for instance, by exciting the atoms to the  $p$ -band of a conventional optical lattice<sup>766–769</sup> or by optically transferring OAM<sup>770</sup> to atoms confined to an arrangement of ring-shaped potentials, which can be created by a variety of techniques, as discussed in Sec. III. Three independent tunneling amplitudes<sup>771</sup> exist in the system:  $J_1$ , which corresponds to the self-coupling at each site between the two OAM states with different circulations, and  $J_2$  and  $J_3$ , which correspond to the cross-coupling tunneling amplitudes between OAM states in different sites with equal or different circulations, respectively. For  $\Theta = \pi/2$ ,  $J_1$  and  $J_3$  acquire a relative phase of  $\pi$  along one of the diagonals of the chain and, due to destructive interference between neighboring sites, the self-coupling vanishes everywhere except for the sites at the left edge. The model possesses inversion and chiral symmetry but, due to the two-fold degeneracy, Zak’s phases<sup>772</sup> are ill-defined. Thus, a series of exact mappings are required to fully characterize topologically the system.



**FIG. 69.** Phase diagram for hard-core bosons on the two-leg ladder and  $U_{\perp} = 0$  as a function of flux per plaquette  $\lambda$  and the interchain hopping  $\Omega$ . The boundaries between the Meissner and the Vortex phase are shown by the red and black solid lines, respectively, for the noninteracting and the interacting case. At  $\lambda = \pi$  the thick solid green-line shows the occurrence of the second incommensuration. The three insets in the right panel show intensity plots of  $n(k, \lambda)$  for the three different values of  $\Omega/J = 0.25, 1.25$  and  $2$ . In panels (b) and (c), the system enters the Vortex phase for  $\lambda > \lambda_c$ : the single peak at  $k=0$  splits in two maxima symmetric around  $k=0$  at  $\pm q(\lambda)$ . At large  $\Omega/J = 2$ , panel (a), the system stays always in Meissner phase and in the vicinity of  $\lambda = \pi$ ,  $n(k)$  becomes independent of  $k$  indicating the formation of a fully localized state (thick solid dark-red line).

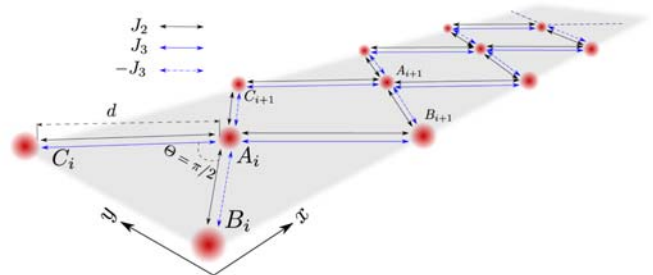


**FIG. 70.** Phase diagram at  $n=0.5$  for a fixed value of interchain hopping  $\Omega/J = 0.5$  as a function of the applied flux  $\lambda$  and as function of the strength of the interchain interaction  $U_{\perp}$ . The dashed black line is the boundary between the Meissner phase and the Meissner-CDW phase (dark-red region). In the Melted vortex phase (light-green region), the dot-dashed line indicates the Lifshitz point. At large  $\lambda$ , the Vortex phase is reestablished (light-blue region under the dotted black line). In the shaded blue area, second-incommensuration occurs. The three insets in the right panel show the behavior of  $n(k)$  (blue solid lines), the spin static structure factor  $S^s(k)$  (solid dark-green line), the charge static structure factor  $S^c(k)$  (solid red line) and the rung current-rung current correlation function  $C(k)$  (solid black lines), for the three points  $A = (\lambda = 0.146\pi, U_{\perp} = -1.5)$ ,  $B = (\lambda = 0.468\pi, U_{\perp} = -1.5)$  and  $C = (\lambda = 0.75\pi, U_{\perp} = -0.25)$  shown in the phase diagram, respectively, for the CDW-Meissner, melted vortex, and vortex phase.

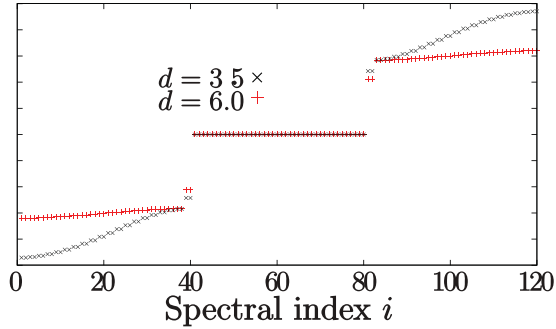
In addition, the model here obtained corresponds to a square-root topological insulator,<sup>773,774</sup> i.e., the quantized values of the Zak's phases are recovered after taking the square of the bulk Hamiltonian.

Under periodic boundary conditions, the diagonalization of the bulk Hamiltonian yields six energy bands in three degenerate pairs and a gap appears in the spectrum. In the  $J_2 = J_3$  limit, all bands become flat. Exact diagonalization, in the case of open boundary conditions, shows the presence of four in-gap states localized at the right edge of the chain, which persist as long as the energy gap is open (see Fig. 72).

We perform first a rotation into a basis of symmetric and anti-symmetric states, which decouples the diamond chain with six states per unit cell into two independent and identical diamond subchains with three states per unit cell. This explains the two-fold degeneracy of the spectrum and the presence of gaps in the band structure. A second basis rotation maps each of the diamond subchains into a modified Su-Schrieffer-Heeger (SSH) model<sup>775</sup> with an extra dangling state per unit cell, which allows us to understand the existence of in-gap edge states localized at the right edge of the chain [Fig. 73(a)], the zero-energy flat band states without population in the central sites



**FIG. 71.** Schematic representation of the considered diamond chain.



**Fig. 72.** Exact diagonalization spectra of a diamond chain of  $N_c = 20$  unit cells for  $d = 3.5\sigma$ , corresponding to  $J_3/J_2 = 1.67$  (black solid line) and  $d = 6\sigma$  corresponding to  $J_3/J_2 = 1.13$  (red dotted line), where  $\sigma = \sqrt{\hbar/(m\omega)}$ .

[Fig. 73(b)] and the flattening of the bands in the  $J_2 = J_3$  limit. Figure 73(c) shows the two degenerate ground states of the system. The decoupled subchains do not have inversion symmetry so that the Zak's phase can yield nonquantized values. Thus, a third mapping to recover inversion symmetry has been introduced<sup>776</sup> obtaining a diamond chain with alternating tunneling amplitudes topologically characterized in Refs. 777 and 778. A striking feature of the topology of this model, directly carried over to the original OAM  $l = 1$  model, is that there is no topological transition across the gap closing point, as can be seen by fixing either  $J_2$  or  $J_3$  and varying the other across zero.

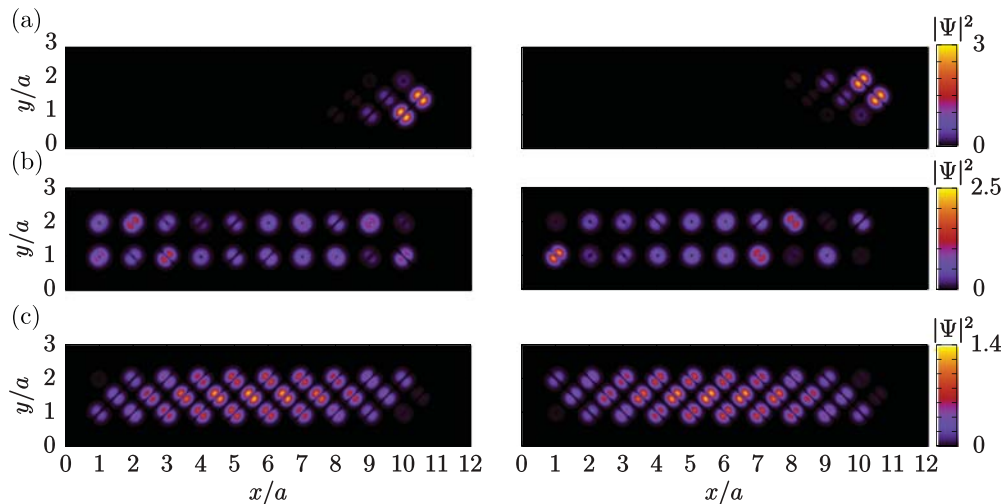
Finally, we have also demonstrated that the system can exhibit Aharonov–Bohm caging in the  $J_2 = J_3$  limit since, in this limit, the states involving the central site of a unit cell can be expressed in terms of flat-band states that occupy solely the four sites surrounding it. Thus, an initial state prepared in an arbitrary superposition of the central sites states will oscillate coherently to its four neighboring sites with a frequency given by the absolute value of the energies of the top/bottom flat-band states without leaving the cage formed by two consecutive unit cells.

## 2. Simulating quantum magnetism with strongly interacting ultracold bosons

Up to here, we have neglected interactions among the ultracold atoms. However, as discussed in Ref. 754 and 755, strongly interacting ultracold bosons loaded into OAM states of lattices of side-coupled cylindrically symmetric traps, e.g., a quasi-one-dimensional ladder of ring potentials or a diamond chain, can realize a variety of spin  $\frac{1}{2}$  models, including the XYZ Heisenberg model with or without external fields. In Ref. 755, we have focused on the Mott insulator regime at unit filling, where each trap is occupied by a single boson and a direct mapping between the degree of freedom corresponding to the two opposite circulations  $\pm l$  of the OAM states to a spin  $\frac{1}{2}$  can be performed. Thus, by tuning the relative phases in the tunneling amplitudes, which depend on the relative orientation between the traps, the system can be used to simulate different spin  $\frac{1}{2}$  models of quantum magnetism. To this aim, we have first computed the explicit dependence of the effective tunneling couplings on the relative angle between the traps by means of second-order perturbation theory. Then, we have discussed for which particular geometries the XYZ Heisenberg model with uniform or staggered external fields could be obtained. As an example, for a quasi-one dimensional ladder of ring potentials with central angle tuned to  $\Theta^l = (2s + 1)\pi/(2l)$  with  $s \in \mathbb{N}$ , single spin flips mediated by interactions do not take place and only isotropic two-spin flips occur. In this situation, the effective Hamiltonian of the system becomes a XYZ Heisenberg Hamiltonian without the external field.<sup>755</sup>

$$H_{\text{eff}}^l = \sum_{j=1}^N J_{xx}^l \sigma_j^x \sigma_{j+1}^x + J_{yy}^l \sigma_j^y \sigma_{j+1}^y + J_{zz}^l \sigma_j^z \sigma_{j+1}^z, \quad (127)$$

where  $J_{xx}^l = -((J_2^l)^2 + (J_3^l)^2)/(2U)$ ,  $J_{yy}^l = -((J_2^l)^2 - (J_3^l)^2)/(2U)$ , and  $J_{zz}^l = -3((J_2^l)^2 - (J_3^l)^2)/(2U)$ , in which  $J_2^l$  and  $J_3^l$  are the cross-coupling tunneling amplitudes between states of OAM  $l$  possessing equal and different circulations, respectively, while  $U$  is the nonlinear interaction parameter.



**Fig. 73.** Density profiles of numerically obtained eigenstates for a diamond chain of  $N_c = 10$  units cells and intersite separation  $d = 6\sigma$ , corresponding to  $J_3/J_2 = 1.13$ . (a) Two degenerate edge states. (b) Two states of the flat band. (c) The two degenerate ground states of the system.

Worth highlighting that besides engineering different spin 1/2 models by tuning the geometry of the lattice, the system also allows one to adjust the relative strength between the effective couplings by changing the radius of the ring traps and their separation. In fact, we have shown that this additional parameter of control can be exploited in realistic experimental setups to explore distinct phases of the XYZ model without external field. Moreover, we have analyzed the effect of experimental imperfections, such as the influence on the tunneling phases of the presence of small fluctuations in the relative angle between the traps. Regarding the physical implementation of the proposal, we have discussed several possibilities to realize a lattice of ring potentials with a tunable geometry and have analyzed single-site addressing techniques that could allow to retrieve the state of each individual spin. Finally, we have also investigated the collisional stability of the system and concluded that the anharmonic energy spacing between OAM states introduced by the ring geometry allows extending the lifetime of the Mott state.

## D. Concluding remarks and outlook

The examples detailed in this section show that the ring geometry allows us to study both the phase diagram and the main features of the excitation spectrum of the infinite ladder to large accuracy as well as to highlight interesting parity and commensurability effects typical of finite rings. Furthermore, the ring geometry allows for new probes of the various phases, e.g., by the measurement of persistent currents or via spiral interferometry. It also displays Josephson modes.

In outlook, one should develop suitable theoretical methods to describe the crossover from the weak-interaction and large occupancy regime down to the strongly correlated regime reached at large interactions and small filling.<sup>779</sup> To make contact with a real experimental situation, it is necessary to investigate how much the signatures of these phases are robust against the finite temperature effects together with the possibility of having long-ranged interactions between the atoms. Also, the experimental realization of ring ladders seems close to reach and would provide a benchmark of atomtronic devices.

## ACKNOWLEDGMENTS

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## XIV. QUANTUM-ENHANCED ATOMTRONICS WITH BRIGHT SOLITONS

**P. Naldesi, J. Polo, S. A. Gardiner, M. Olshanii, A. Minguzzi, L. Amico**

Quantum coherent states of macroscopic degrees of freedom are hard to achieve, due to decoherence. Attractive bosons are a very special case study, completely different from repulsive bosons. In this

chapter we revise their properties in the quantum regime, such as the chemical composition of its ground state, the transmission across a barrier, the excitation spectrum and the response to rotation, showing that they provide a new type of resource for atomtronics applications.

## A. Scattering properties of attractive bosons against a barrier

Cold bosonic systems tend to remain in a Bose-condensed state that can be perfectly described by a mean-field theory. However, for attractive condensates, there are points in the space of parameters where the mean-field theory predicts sudden jumps for some macroscopic observables. The relevant example corresponds to a single one-dimensional bosonic soliton<sup>780–782</sup> or any other many-body bound state that is scattered off a barrier in a typical scattering setup, where a localized wavepacket is prepared and sent toward an obstacle.<sup>783–785</sup> In such scattering events, the nonlinearity of the mean field theory can lead to some “unsettling” results. For instance, for incident kinetic energies (per particle) below  $\frac{1}{4}$  of the magnitude of the soliton chemical potential, a “forbidden window,” in the form of a discontinuity, on the axis of the transmission coefficients must emerge<sup>786</sup> (see also Refs. 547 and 787); it appears due to the fact that the amount of the incident kinetic energy is insufficient to compensate for the loss of the interaction energy in a 50%–50% splitting in this regime.

As we scan the barrier height from a lower value up, the transmission coefficient increases and, at some barrier height, abruptly jumps up.<sup>786</sup> At the mean-field level, the jump is infinitely sharp. Indeed, a dissociation of the soliton onto the transmitted and reflected parts costs interaction energy, and the incident kinetic energy may not be sufficient to pay for it (see Fig. 74).

Such a discontinuity is nonphysical. As has been shown in Refs. 549 and 788, the key to ensure the continuity of the transmission coefficient curve is to recognize that at the apparent discontinuity point, the condensate becomes fragmented and the transmission events acquire a quantum randomness. This regime will soon be within experimental reach.<sup>784</sup> The good news is that a highly desirable Schrödinger cat is itself a fragmented state; the bad news is that if the number of occupied one-body orbitals becomes large, the macroscopic coherence becomes unusable. Reference<sup>532</sup> suggests a secure way of suppressing the undesirable fragments: the soliton kinetic energy must be decreased even further to a point where the *total* kinetic energy becomes less than the chemical potential, thus ensuring no relative motion of the constituent atoms with only “cold soliton transmitted” and “cold soliton reflected” allowed orbitals as the result. While conceptually elegant, this method of generating a macroscopic coherence requires center-of-mass kinetic energies  $N$  times lower than those currently used ( $N$  being the number of atoms in the soliton) and

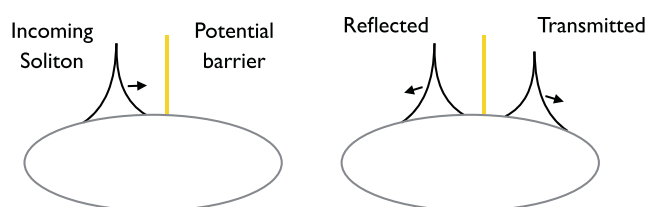


Fig. 74. Scheme of a soliton scattered from a potential barrier.



scattering regimes where the barrier becomes completely classical from the soliton center-of-mass point of view. Accordingly, Ref. 549 suggests using extended center-of-mass coherent wavepackets with a nonzero velocity width, where the barrier is used as a classical velocity filter—see also Ref. 545. However, a private communication,<sup>789</sup> and by careful inspection of Figs. 2 and 3 in Ref. 788, indicates—based on numerical evidence—that even at moderate kinetic energies, there remain only two populated orbitals. If this is indeed the case, then it is clear what these orbitals are: they are nothing else but the state of the condensate just before and just after the mean-field jump in the transmission coefficient.

Note that even in the favorable two-orbital case, the macroscopic coherence may still remain unusable due to the entanglement between the center-of-mass motion and possible excitations created during the scattering event. Even if these excitations are small at the level of the BEC wavefunction, the difference between the internal states of the transmitted and reflected condensates may still be large due to the orthogonality catastrophe. Nonetheless, the macroscopic coherence may be potentially preserved if a limited number of atoms is used. The upper bound for this number is yet an open question, which will require an intensive numerical study.

While in the proposal,<sup>549,788</sup> the center-of-mass of the incident soliton is assumed to be in a coherent state prior to splitting, the cooling of a macroscopic variable to that state is difficult *per se*. However in Refs. 790 and 791, it is shown that in a factor of four quench of the coupling constant, one can create, while at a finite temperature, an exponentially cold quantum state of a relative distance between the centers-of-mass of two solitons<sup>792–795</sup> itself a macroscopic variable. The theoretical estimates<sup>791</sup> show that under realistic experimental conditions, quantum fluctuations of the intersoliton velocity will lead to an observable intersoliton separation after a time

$$\tau \approx 4.7 \text{ s.}$$

Further beam-splitting of the intersoliton distance degree of freedom requires additional study, while its initial coherence is already guaranteed. Classically, the above states would correspond to Gross–Pitaevskii breathers<sup>796,797</sup> with fluctuating parameters. These have recently been experimentally realized, albeit in the classical regime, as described in Ref. 798.

## B. Creation and manipulation of quantum solitons

### 1. Quantum solitons in the Bose–Hubbard model

Attractive bosons confined in a one dimensional lattice system can be described by the Bose–Hubbard model. Before moving to atomtronics applications, e.g., a ring lattice, we present here the properties of the ground state and its excitations. For a mesoscopic sample, with the limited number of bosons, higher band occupancies are negligible<sup>721</sup> and, even at intermediate and strong attractions, the occupancies can be integrated in the model by renormalizing the tunneling and interaction parameters.<sup>799–801</sup> In this regime, the Hamiltonian reads

$$\hat{\mathcal{H}} = -J \sum_{j=1}^L \left( a_j^\dagger a_{j+1} + \text{h.c.} \right) - \frac{|U|}{2} \sum_{j=1}^L \hat{n}_j (\hat{n}_j - 1), \quad (128)$$

where the operators  $a_i^\dagger$  obey the canonical commutation relations  $[a_i, a_j^\dagger] = \delta_{ij}$ ,  $n_i = a_i^\dagger a_i$  is the number operator at the site  $i$ , and the

operators  $a_i$  and  $L$  is the number of sites in the chain. The parameters  $J$  and  $U$  in Eq. (128) are the hopping amplitude and the strength of the on-site interaction, respectively. Periodic boundary conditions are implemented, requiring that  $a_1^\dagger a_L = a_L^\dagger a_1$ . The lattice is loaded with  $N$  bosons. While some exact results are available for  $N = 2$ <sup>711,802,803</sup> and an effective model can be built to explain the spectrum for  $N = 3$ ,<sup>804,805</sup> for a larger number of particles, the system is not solvable and numerical simulations are necessary.<sup>806,807</sup>

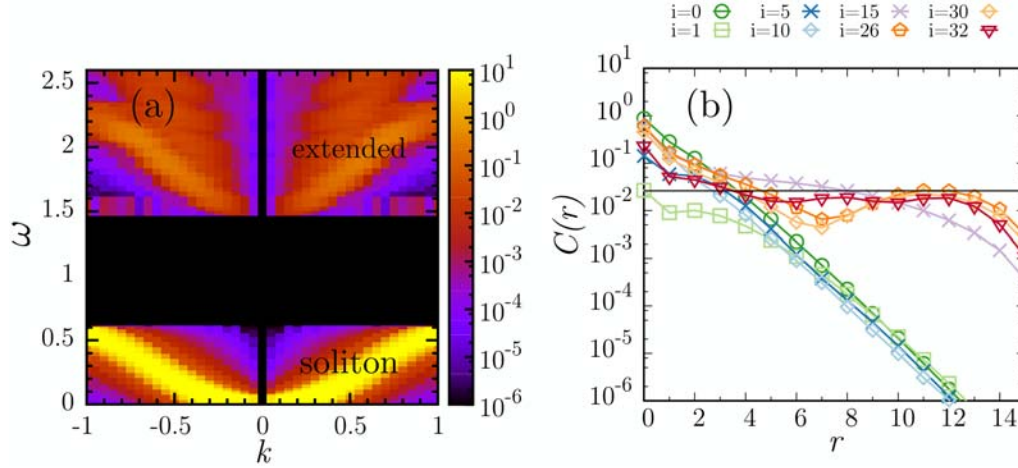
*a. 2-particle sector.* The problem of two attracting bosons on a lattice is exactly solvable à la coordinate Bethe ansatz by transforming the wave function in the center of mass and relative coordinates. This solution is also valid in the presence of an synthetic gauge field.<sup>803</sup> The eigenstates of the system form two bands depending on the nature of relative momentum. For imaginary solutions, we have the lowest energy branch composed by the  $L$  bound-state (solitons), while the real solutions correspond to scattering states that form the second band at higher energy. The energy gap separating the two increases with interactions, and it is found that for  $U/J \geq 4$ , the two bands completely detach for each momentum.<sup>803</sup>

*b. N-particle sector.* For a larger number of particles, the BHM [Eq. (128)] is not solvable by the coordinate Bethe ansatz. The failure results because of finite probabilities that a given site is occupied by more than two particles,<sup>808–810</sup> whose interaction cannot be factorized in two-body scattering.

Information on the available excitations in the system as a function of their momentum  $k$  and energy  $\omega$  is provided by the dynamical structure factor  $S(k, \omega)$ ,

$$S(k, \omega) = \sum_{\alpha \neq 0} \sum_r |\langle \alpha | e^{-ikr} \hat{n}_r | 0 \rangle|^2 \delta(\omega - \omega_\alpha). \quad (129)$$

where  $\hat{n}_r$  is the number operator acting on the site  $r$  and  $|0\rangle$  is the ground state and  $\alpha$  labels the states with increasing energy (i.e.,  $\alpha = 1$  is the first excited state). The peaks of  $S(k, \omega)$  reconstruct the energy bands of the system<sup>804,805</sup> and are shown in Fig. 75. Numerical results show a scenario similar to the two-particle case with a low-energy band that is separated from the rest of the spectrum. The nature of such a band can be analyzed by the study of correlation functions,  $C(r) = \langle n_{L/2} n_{L/2+r} \rangle$ . The numerical analysis shows that the lowest energy band is composed of many-body bound states. In fact, all these states are characterized by an exponential decaying of correlations  $C(r) \sim \exp(-r/\xi)$ . The correlation length  $\xi$  is fixed only by the interactions and decreases with increasing  $U$ . For states belonging to the second branch  $C(r)$  approaches, at intermediate distances, a plateau  $\sim n_{as} = (N/L)^2$ , before dropping down when approaching the walls of the box. We, thus, can conclude that the higher branch contains extended states. Notice that, at difference from the continuum case, where a Bethe ansatz solution is available and one can tell the nature of the state by checking whether the rapidities are real or complex in the lattice case, there is no exact solution; hence, no way to tell whether they are scattering states,  $N - 1$ -body bound states, etc. Thus, the dynamical structure factor is very practical to visualize all types of excitations. Also, in this case, the bands gap increases with interactions and the critical interaction to have a complete detachment of the bands scales like  $U_c \sim 1/N$ .



**FIG. 75.** Panel (a): dynamical structure factor  $S(k, \omega)$  for a chain of  $L = 30$  sites. Numerical results for  $N = 5$  particles and interactions  $U = 1.2$ . Panel (b): Density-density correlation function  $C(r)$  for  $N = 5$  particles in a chain of  $L = 30$  sites and interactions  $U = 0.6 < U_c$ . Correlations are computed over several excited states labeled by  $i$  ( $i$ -th excited state,  $i = 0$  correspond to the ground state).

*c. Soliton stability.* Finally, we devise a specific dynamical protocol to study the soliton stability and evidence the features of the band structure.

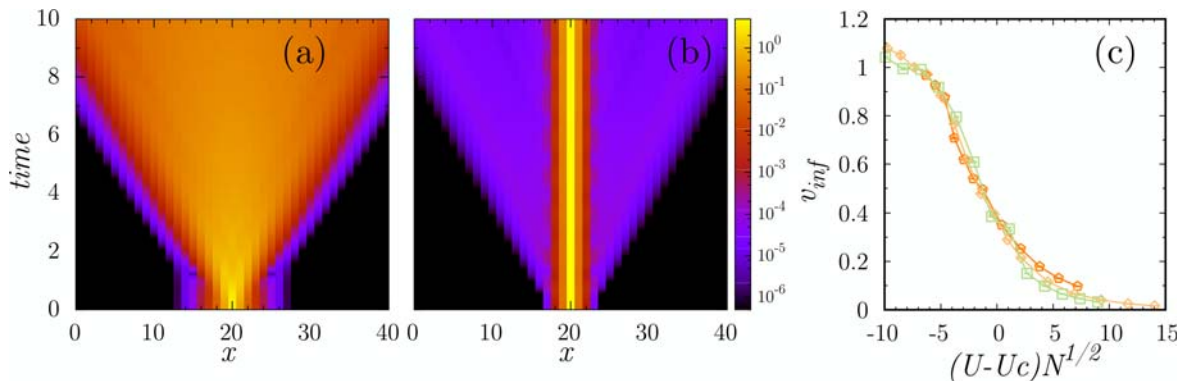
By initially breaking the lattice translational symmetry with an attractive potential  $\mathcal{H}_i(\mu, U) = \mathcal{H}(U) + \mu(U)n_{i_0}$ , a soliton is pinned in a given site  $i_0$  of the lattice, and then, let it expand by removing the pinning. In this way, while for small  $U$ , we populate both scattering and bound states for  $U > U_c$ , when the gap separates the two bands, mostly bound states are populated.

In Figs. 76(a) and 76(b), we show the expansion dynamics of the density for two cases:  $U < U_c$  and  $U > U_c$ . Increasing the interaction strength, the density profile remains closer and closer to the one of the initial state. Only a small fraction is spreading into the chain leading to a higher stability of the soliton. This phenomenon can be studied more quantitatively by analyzing the expansion velocity,  $v(t) = (d/dt)\sqrt{R^2(t) - R^2(0)}$ , with  $R^2(t) = (1/N)\sum_{i=1}^L n_i(t)(i - i_0)^2$  and its asymptotic value at large times  $v_\infty$ . The inspection of  $v_\infty$  in Fig. 76(c) further shows the difference

between the two regimes. While there is no criticality in the system close to  $U_c$ ,  $v_\infty$  displays a peculiar scaling behavior and acts like an order parameters for the system.

## 2. Solitons in rotation

As an application to atomtronics, attracting bosons can be used to devise a new type of interferometer, based on superposition of persistent current states. The effect of an induced rotation, or more generally of a (synthetic) gauge field, on such a system has been extensively studied.<sup>591</sup> It is in fact well known that the quantum system in a ring geometry displays a staircase response to an applied gauge field of intensity  $\Omega$ . The induced angular momentum increases in quantized steps as a function of  $\Omega$ ,<sup>142,432</sup> and the amplitude of persistent currents displays periodic oscillations with  $\Omega$ .<sup>811,812</sup> The periodicity of such oscillation is completely fixed by the effective flux quantum present in the system and does not depend on the intensity of particle-particle interactions.<sup>813</sup> In the following, without loss of generality, we will



**FIG. 76.** Panels (a) and (b): expansion of a soliton composed by  $N = 5$  particles, pinned to the center of a chain with  $L = 41$  sites for interactions  $U = 0.4$  and  $U = 1.8$ . Panel (c): asymptotic expansion velocity  $v_\infty$  as a function of  $U - U_c(N)$  and of  $(U - U_c(N))\sqrt{N}$ .

refer only to the case of an artificial gauge field induced by a global rotation at angular frequency  $\Omega$ . Our discussions can be applied to any type of artificial gauge fields.

For strongly correlated one-dimensional bosons with attractive interactions, as we discuss in the following, the nature of flux quantum is nontrivial due to the formation of many-body bound states. This feature has dramatic effects on the persistent current that oscillates with a periodicity  $N$  times smaller than in the standard case corresponding to repulsive interactions. Remarkably, the periodicity depends on the interaction, which leads to an extension of the Leggett theorem.

*a. Continuous ring.* For a continuous ring, the system can be described through the Bose-gas integrable theory, i.e., the Lieb–Liniger model.<sup>814</sup> This is the case when the density  $N/L$  of bosons, where  $N$  is the particle number and  $L = 2\pi R$  is the perimeter of the ring of radius  $R$ , is small. A well established limiting procedure allows one to link the lattice and continuous models (see, e.g.,<sup>803</sup> and <sup>814</sup> for a discussion). For such systems, exact results are well known.<sup>251</sup> The Lieb–Liniger Hamiltonian in the rotating frame reads

$$\hat{\mathcal{H}}_{LL} = \sum_{j=1}^N \frac{1}{2m} (p_j - m\Omega R)^2 + g \sum_{j<l} \delta(x_j - x_l) - E_\Omega, \quad (130)$$

where  $m$  and  $p_j$ 's are the mass and the momentum of each particle, respectively;  $L_z = \sum_{j=1}^N L_{z,j}$  is the total angular momentum of the  $N$  particles;  $g$  is the interaction strength; and  $E_\Omega = Nm\Omega^2 R^2/2$ .

The solution of the model dramatically change according to the sign of the interactions. For repulsive interactions, independently on their strength, the ground state energy  $E_{GS}$  is periodic in  $\Omega$  with period  $\Omega_0 = \hbar/mR^2$ . The persistent current in the rotating frame defined as  $I_p = -(\Omega_0/\hbar)\partial E_{GS}/\partial\Omega$  displays a sawtooth behavior versus  $\Omega$ ,<sup>813</sup> corresponding to a staircase behavior of angular momentum  $L_z$ .

For attractive interactions, the scenario changes completely; the ground state is a many-body bound state, i.e., a “molecule” made of  $N$  bosons, corresponding to the quantum analog of a bright soliton.<sup>806,815,816</sup> The ground state energy for arbitrary  $\Omega$  then reads

$$E_{GS} = \frac{\hbar^2}{2MR^2} \left( \ell - N \frac{\Omega}{\Omega_0} \right)^2 - \frac{N(N^2 - 1)g^2}{12}, \quad (131)$$

where the second term accounts for the interaction energy  $E_{int}$  and is independent on the rotation frequency. This result clearly shows how, under the effect of the artificial gauge field, attracting bosons effectively behave as a single massive object of mass  $M = Nm$ . The energy displays a  $1/N$ -periodicity as a function of the artificial gauge field,  $\Omega$ , in units of  $\Omega_0$  corresponding to *fractionalization* of angular momentum per particle.

*b. Lattice ring.* When the density of particles is not small, the lattice effects that break the integrability of the model start to be relevant. In this situation, the system is well described by the Bose–Hubbard model (BHM),

$$\hat{\mathcal{H}}_{BH} = \sum_{j=1}^{N_s} \frac{U}{2} n_j(n_j - 1) - J \left( e^{-i\hat{\Omega}} a_j^\dagger a_{j+1} + \text{h.c.} \right), \quad (132)$$

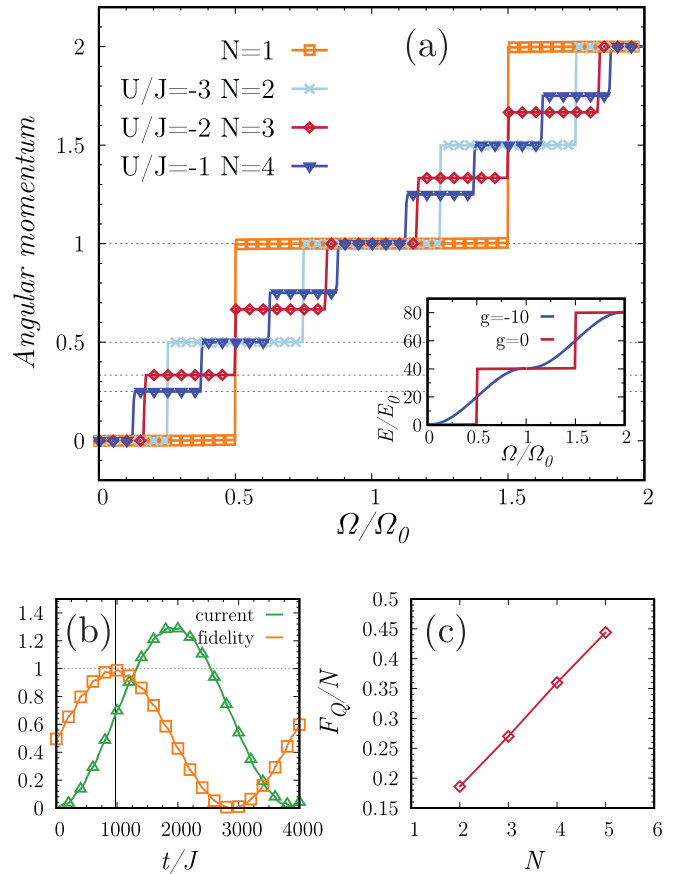
where  $a_j$  and  $a_j^\dagger$  are site  $j$  annihilation and creation Bose operators and  $n_j = a_j^\dagger a_j$  is the number operator. The parameters  $J$ ,  $U < 0$  in Eq.

(132) are the hopping amplitude and the strength of the on-site interaction, respectively;  $N_s$  being the number of sites in the lattice and  $\Omega \doteq 2\pi\Omega/(\Omega_0 N_s)$  for brevity.

In the lattice model [Eq. (132)], the center-of-mass and relative coordinates, at any finite interaction, cannot be decouple. This feature has a profound implication on the behavior of persistent current. As we will discuss below, in contrast with the continuous theory, here the persistent current periodicity does depend on interaction strength.

In Fig. 77, we show the numerical results of angular momentum: also, in this case, the  $1/N$  periodicity in  $\Omega/\Omega_0$  of the persistent currents as well as the fractionalization of angular momentum emerges. While fractionalization always occurs, the  $1/N$  periodicity is affected by the interplay between the system size and interaction strength.

When interactions are sufficiently large, the “size of the many-body bound state,” i.e., the decay length of the density–density correlations,<sup>806</sup> is much smaller than the size of the system. Upon decreasing the interactions, the size of the many-body bound state increases more and more over the chain and the solitonic nature of the state gets less



**Fig. 77.** Panel (a): average angular momentum per particle (inset: GP analysis) as a function of the artificial gauge field for different particle number and for particular values of interaction strength. Panel (b): time dependent current (in units of the hopping constant  $J$ ) following a quench from  $\Omega/\Omega_0 = 0$  to  $\Omega/\Omega_0 = 1/2$ . Here we set  $L = 28$ ,  $N = 3$ ,  $U/J = -0.51$  and  $\Delta_0/J = 0.015$ . Panel (c): Quantum Fisher information as a function of the particle number showing the Heisenberg-limited behavior  $F_Q \propto N^2$ .

and less pronounced. All the observed features are purely quantum many-body effects tracing back to specific quantum correlations since they completely disappear in a mean-field Gross-Pitaevskii description of the system.

Angular momentum fractionalization and the related persistent current periodicity can be observed with standard time-of-flight (TOF) techniques. Measuring the distributions of the atoms after releasing the trap confinement and turning off interactions, we have access to the momentum distribution defined as  $n(\mathbf{k}) = |w(\mathbf{k})|^2 \sum_{j,l} e^{i\mathbf{k} \cdot (\mathbf{x}_j - \mathbf{x}_l)} \langle a_j^\dagger a_l \rangle$ . In fact, we find that the mean-square radius of the distribution increases in fractional steps of for  $\Omega/\Omega_0 = \ell/N$ .<sup>126</sup>

### 3. Entangling solitons with different $L_z$

We finally demonstrate how the scenario above can be harnessed to create specific entangled states of persistent currents. Such entangled states are characterized by an increased sensitivity to the effective magnetic field that reaches the Heiseberg limit. In the following, we propose a specific dynamical protocol that allows us to create such type of state.

Since the Hamiltonians in Eqs. (130) and (132) commute with the total angular momentum, dynamically mix entangle states with different angular momentum, the rotational invariance of the system needs to be broken. The ring is then interrupted with a potential barrier of strength  $\Delta_0$  localized in a single lattice site. Then, the artificial gauge field is quenched from  $\Omega = 0$  to  $\Omega = \Omega_0/2$ . This procedure is capable of dynamically entangling the angular momentum state at  $\Omega = 0$ , i.e.,  $L_z = 0$ , with the one at  $\Omega = \Omega_0$ , i.e.,  $L_z = N$  (see Fig. 77), yielding  $|\psi\rangle_{\text{NOON}} = \frac{1}{\sqrt{2}}(|L_z = 0\rangle + |L_z = N\rangle)$  when the current reaches the half of its maximum value.

The response of such a state to an external rotation is  $|\psi(\phi)\rangle = e^{i\phi \hat{L}_z/\hbar} |\psi\rangle_{\text{NOON}}$ , and the quantum Fisher information<sup>817,818</sup>  $F_{\mathcal{Q}} = 4\langle (\psi'(\phi)|\psi'(\phi)) - |\langle \psi'(\phi)|\psi(\phi) \rangle|^2 \rangle$ , being  $\psi'(\phi) = \partial|\psi(\phi)\rangle/\partial\phi$ . For our state, we find  $F_{\mathcal{Q}} \sim N^2$ , i.e., it reaches the Heisenberg limit—see Fig. 77. The corresponding sensitivity  $\delta\phi$ , therefore, is

$$\delta\phi \geq \frac{1}{(F_{\mathcal{Q}})^{1/2}} = \frac{1}{N}. \quad (133)$$

This shows that entangled states of quantum solitons with different angular momenta lead to a quantum advantage of the sensitivity for rotation detection. Notice that this type of entangled state is completely different from a superposition state obtained by splitting a soliton by a barrier in real space, which could be used, e.g., as a gravimeter. In both cases, the main experimental limitation is due to unwanted fluctuations (thermal, technical, etc.) and particle losses. The latter are nevertheless expected to play a minor role for the small particle numbers considered in this setup.

### C. Concluding remarks and outlook

In this section, we studied attractive bosons in the quantum regime. Its ground state is an  $N$ -body bound state, which on a lattice is protected by a gap with respect to the first branch of excitations, corresponding to scattering states. We have shown that this implies the stability of a soliton initially prepared in a pinning site. We have also shown that quantum solitons on a ring display an enhanced response

to artificial gauge field  $\Omega$  with a  $1/N$  periodicity as a function of  $\Omega/\Omega_0$ . This corresponds to fractionalization of angular momentum per particle, intrinsically due to the presence of many-body bound states. Finally, we have identified a protocol to create a nonclassical superposition of angular momentum states by a suitable quench of the artificial gauge field based on angular momentum fractionalization. The use of quantum coherent macroscopic superposition states in atom interferometry devices can increase considerably the phase sensitivity. The states studied in this section can yield an  $N$ -fold enhancement in sensitivity to rotation in a ring-based gyroscope. In a typical configuration, a localized barrier can split the solitons in two waves propagating in clockwise and anticlockwise that can ultimately recombine producing interference fringes with a specific pattern. Controlling the effects of decoherence, losses, and the identification of the optimal working parameters for bright solitons based interferometers is important challenges to overcome.

## XV. ATOMTRONICS WITH ALKALINE-EARTH-LIKE METAL ATOMS

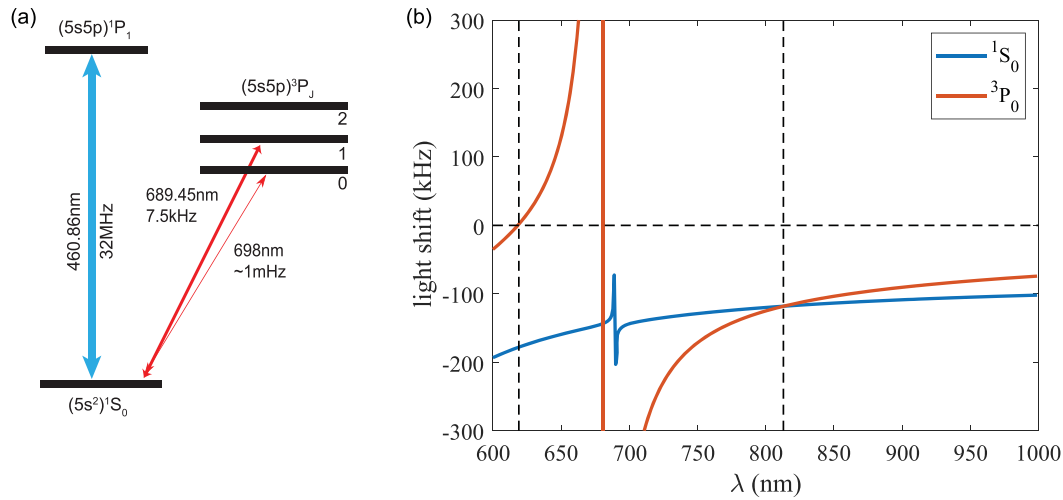
**D. Wilkowski, W. J. Chetcuti, C. Miniatura, L.-C. Kwek, L. Amico**

### A. Why alkaline-earth-like metal atoms?

Over the past two decades, the number of experiments using ultracold alkaline-earth-like metal atoms have considerably increased. Indeed, these atoms have singlet and triplet electronic spectra that offer interesting alternatives over the usual doublet spectrum of the more commonly used alkali metal atoms. For the purpose of illustration, we show the energy levels and transitions of interest for strontium atoms (Sr) in Fig. 78(a). Laser cooling and magneto-optical traps are achieved using the electric dipole-allowed singlet  $^1S_0 \rightarrow ^1P_1$  transition. For heavy elements (Sr, Yb, Hg), the singlet-triplet intercombination line is strong enough to allow further cooling. For example, for Sr atoms, reaching temperatures at the single photon recoil limit on large atomic ensembles<sup>819</sup> can be achieved simply with the usual Doppler cooling technique.<sup>820</sup> In addition, since multiple scattering is limited, large space phase densities can be reached compared to alkali metal atoms.<sup>821</sup> Laser cooling on intercombination lines is then efficient and provides a favorable starting point to reach quantum degeneracy with evaporative cooling techniques. The latter was obtained for several isotopes of Yb and Sr, such as  $^{84}\text{Sr}$  (0.6%),<sup>822,823</sup>  $^{86}\text{Sr}$  (9.9%),<sup>824</sup>  $^{87}\text{Sr}$  (7.0%),<sup>825,826</sup>  $^{174}\text{Yb}$  (31.8%),<sup>827</sup>  $^{173}\text{Yb}$  (16.1%),<sup>828</sup>  $^{176}\text{Yb}$  (12.8%),<sup>829</sup> and also for  $^{40}\text{Ca}$  (96.9%).<sup>830</sup> The percentage, given in parentheses, is the relative abundance of the isotope.

Importantly, we note that the spin-singlet ground state is not sensitive, or only weakly so (for nuclear spin of fermionic isotopes), to magnetic fields. Magnetic trapping is, thus, excluded as well as the possibility of using magnetic Feshbach resonances to tune the scattering length and, in turn, interactions. One has to rely on optical dipole traps and zero-field interactions to implement evaporative cooling. There have been attempts to control the scattering length by optically dressing the ground state level to some molecular bound states in the excited level<sup>831</sup> but the lifetimes of such dressed states remain too short to be of practical interest.<sup>832</sup>





**FIG. 78.** (a) Energy levels and transitions of interest for Strontium. The transitions linewidths and wavelengths are indicated in the figure. (b) Light shifts of the  $^1S_0$  and  $^3P_0$  states of Sr as a function of the wavelength of the dipole trap. The excited-state and ground state light shifts are exactly equal at the magic wavelength 813 nm. The light shifts are calculated for a laser power of 600 mW and a beam waist of 65  $\mu\text{m}$ .

In addition to the electric-dipole transition and the intercombination lines, alkaline-earth-like metal atoms possess a clock transition connecting the ground state  $^1S_0$  to the long-lived excited state  $^3P_0$ . Since these states are energetically well separated, their light shifts, induced by a far-off-resonant laser light, can be engineered almost with high control. An illustrative example is given in Fig. 78(b) for Sr. For instance, in the so-called magic configuration, where light shifts exactly compensate, the transition frequency becomes almost insensitive to the trapping optical field leading to applications in precision frequency and time measurements.<sup>833</sup> Here, atoms are trapped in optical lattices to allow for long interrogation times in a massively parallel configuration, giving the best clock uncertainty to date.<sup>834</sup> Aside from obvious metrological applications, the clock transitions are also suitable for strongly correlated many-body phenomena that may be difficult to be addressed elsewhere, such as the Kondo effect and the heavy Fermion manifestation.<sup>835,836</sup> These experiments can be performed in an optical lattice with a wavelength corresponding to weak light shifts in the excited state and a stronger confinement in the ground state [ $\sim 619$  nm for Sr as shown in Fig. 78(b)]. Moreover, the fermionic isotopes (Sr and Yb) possess a nonzero nuclear spin. Thus, a spin exchange interaction between two atoms is present in the cold collision regime: one in the fundamental orbital  $^1S_0$  and the other in the excited orbital  $^3P_0$ .<sup>837</sup> The resonant character of this Feshbach-type exchange interaction has been shown for Yb<sup>838,839</sup> and opens the door for quantum simulation of strongly correlated 2-orbital quantum gases.<sup>836</sup>

The fermionic isotopes,  $^{87}\text{Sr}$  et  $^{173}\text{Yb}$ , have also the interesting property of possessing a nuclear spin larger than  $\frac{1}{2}$  ( $I = 9/2$  and  $I = 5/2$ , respectively,) decoupled from the electronic orbital<sup>840,841</sup> (in addition to the above-mentioned spin exchange interaction). As a consequence, the many-body Hamiltonian of these systems do not depend on the nuclear spin orientation: They are invariant under the  $\text{SU}(N)$  symmetry with a dimension  $N = 2I + 1$  much larger than  $N = 2$  (corresponding to a spin-1/2 fermion), going thus beyond the usual  $\text{SU}(2)$  symmetry. Numerous theoretical efforts have been pursued to better

understand such  $\text{SU}(N)$  systems, in particular, their magnetic<sup>840</sup> and topological properties and their quantum phase transitions.<sup>842</sup> On the experimental side, important results have been obtained on the Yb Mott insulator.<sup>843,844</sup> Ordered magnetic phases above the Néel temperature could be observed<sup>845</sup> because the entropy per spin component was reduced by the Pomeranchuk effect, relaxing the temperature constraint on the gas.<sup>840,846</sup>

The Sr fermionic isotope has also been used to generate artificial gauge fields. An effective spin-orbit coupling, mediated by the clock transition has been studied in a one-dimensional lattice.<sup>847,848</sup> The goal here is to act on the ultracold gas to obtain many-body states of metrological interest. In another work, non-Abelian gauge transformations have been reported using two dark states of a tripod laser scheme.<sup>849</sup> This configuration appears to be promising for atomtronics and will be discussed in more detail in Sec. XV B.

## B. Effective Abelian and non-Abelian gauge fields

We discuss here the implementation of effective gauge fields for ultracold alkaline-earth metal atoms in the presence of a general atomtronics circuit. We require the spatial scales of the atomtronics circuit to be larger than the laser wavelength used to create the gauge field such that the adiabatic approximation always holds.<sup>591</sup> State differently, the artificial gauge field should act in continuous bulk space and be unaltered by the presence of the atomtronics circuit. This excludes lattice-type structures but one can still implement the gauge field using off-resonant coherent Raman beams as used by Spielman *et al.*<sup>727</sup> Alternatively,  $\text{SU}(2)$  gauge fields can be generated within the dark-state manifold of a four-level resonant tripod scheme.<sup>726</sup> Using a double tripod scheme, the symmetry can be further extended to  $\text{SU}(3)$ .<sup>850</sup> Since dark states are sensitive to ground state energy fluctuations, this scheme is appropriate for alkaline-earth-like metal fermions, which only possess a nuclear spin well protected from their environment (magnetic fields, collisions).

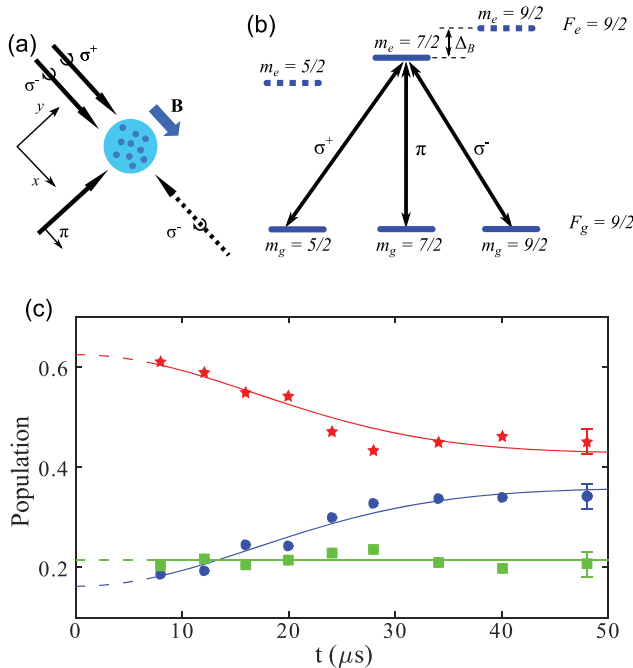
Recently,<sup>849</sup> we implemented a tripod scheme on a cold gas of  $^{87}\text{Sr}$  containing around  $10^5$  atoms using the  $F_g = 9/2 \rightarrow F_e = 9/2$  intercombination line at 689 nm. The cold sample<sup>819</sup> is prepared in a crossed optical dipole trap where atoms are optically pumped in the stretched  $m = F_g$  magnetic substate and Doppler cooled at a temperature around  $0.5 \mu\text{K}$ . A magnetic bias field isolates a particular tripod scheme among the excited and ground Zeeman substate manifolds. The three coupling laser beams are set on resonance with their common  $|m = 7/2, F_e = 9/2\rangle$  excited state [see Figs. 79(a) and 79(b)].

In the adiabatic approximation, the Hamiltonian describing the quantum state evolution in the dark-state manifold<sup>591</sup> reads

$$H = \frac{1}{2M}(\hat{\mathbf{p}}\mathbb{1} - \mathbf{A})^2 + W, \quad (134)$$

where  $\hat{\mathbf{p}} = -i\hbar\nabla$  is the momentum operator,  $\mathbb{1}$  is the identity operator in the internal dark-state subspace, and  $M$  is the atom mass. With equal and constant Rabi frequencies amplitudes, and for the orientation of our laser beams [see Fig. 79(a)], the vector and scalar potential are<sup>849</sup>

$$\mathbf{A} = \frac{2\hbar(\mathbf{k}_2 - \mathbf{k}_1)}{3} \mathcal{M}, \quad W = -\frac{4E_R}{9} \mathcal{M}, \quad (135)$$



**Fig. 79.** (a) Propagation directions of the tripod laser beams (full arrows) and their polarizations along a magnetic field bias ( $B = 67 \text{ G}$ ). For the non-Abelian gauge field discussed in the text, the  $\sigma^-$  laser beam direction is flipped (dashed arrow). (b) Energy levels and experimentally relevant transitions. The magnetic bias field lifts the degeneracy of the different Zeeman manifolds and allows to address each transition individually. The Landé  $g$ -factors are indicated for each hyperfine level. The black arrows connecting the ground state to the excited states correspond to the tripod beams. (c) Time evolution of the bare-state populations after the tripod ignition. Red stars, green squares, and blue circles correspond to  $|m = 9/2\rangle$ ,  $|m = 7/2\rangle$ , and  $|m = 5/2\rangle$  states, respectively. Adapted with permission from Leroux *et al.*, Nat. Commun. **9**, 3580 (2018). Copyright 2018, Author(s) licensed under a Creative Commons Attribution 4.0 License.

where  $E_R = \hbar\omega_R = \hbar^2 k^2 / (2M)$  is the atomic recoil energy and  $\mathbf{k}_j$  is the wavevector of laser beam  $j = 1, 2, 3$  (with  $1 \equiv \sigma^+$ ,  $2 \equiv \pi$ , and  $3 \equiv \sigma^-$ ). The matrix  $\mathcal{M}$  reads

$$\mathcal{M} = \begin{pmatrix} 3/4 & -\sqrt{3}/4 \\ -\sqrt{3}/4 & 1/4 \end{pmatrix}. \quad (136)$$

Since the components of the vector potential commute, this gauge field is Abelian. In Fig. 79(c), we show the result of a ballistic expansion of the cold atomic cloud on the bare state populations. The red stars, green cubes, and blue triangles correspond to the  $|m = 9/2\rangle$ ,  $|m = 7/2\rangle$ , and  $|m = 5/2\rangle$  populations, respectively, whereas the curves correspond to the evolution given by the Hamiltonian, Eq. (134), with thermal averaging.<sup>849</sup> The relaxation of the  $|m = 9/2\rangle$  and  $|m = 5/2\rangle$  populations is due to the thermal averaging, and the temperature is proportional to the characteristic relaxation time of the system.<sup>849</sup>

Remarkably, since the gauge field is Abelian and homogeneous, the field strength (or Berry curvature), i.e., the curl of the vector potential, is zero. Hence, there is no Lorentz forces acting on the system. This result can be simply understood through a simple physical argument: Mechanical forces here come from photon redistribution among the tripod lasers with different propagation directions. With our laser configuration, such photon exchanges would induce a population change only for state  $|m = 7/2\rangle$ . However, this population remains constant, confirming the absence of light-assisted forces.

The situation becomes more complex if one flips the direction of one of the laser along the  $x$ -axis. For instance, if the laser  $\sigma^-$  is flipped, the gauge potential now reads

$$\mathbf{A} = \frac{2\hbar}{3}(\mathbf{k}_1\mathcal{N} + \mathbf{k}_2\mathcal{M}), \quad (137)$$

where

$$\mathcal{N} = \begin{pmatrix} 9/4 & \sqrt{3}/4 \\ \sqrt{3}/4 & 3/4 \end{pmatrix}. \quad (138)$$

The two components of the vector potential do not commute, so the gauge field becomes non-Abelian. In the context of atomtronics, spin precession and related spin-orbit-like coupling can play an important role. For instance, the spin precession leads to spatial oscillation of the wave-packet (as in relativistic *Zitterbewegung* effect), whereas the momentum operator still commutes with the Hamiltonian.<sup>68,851,852</sup> Moreover, it was shown that the characteristic double-well energy dispersion of a spin-orbit coupled system leads to a Josephson effect in momentum space with the presence of supercurrents.<sup>853</sup> Further applications and potential research objectives are given in Sec. XVC.

### C. Persistent current of $\text{SU}(N)$ fermions

Atomtronics can provide key contributions to mesoscopic physics, exploring physical situations that are hard, if not impossible, to explore with standard implementations. One of the purest expressions of mesoscopic behavior is the persistent current. There have been several studies on the persistent current of bosonic systems in ring-shaped circuits. In this article, these are summarized in Secs. VII and IX. Atomtronic circuits comprised of ultracold fermions are much less explored. In Ref. 854, the persistent current of interacting

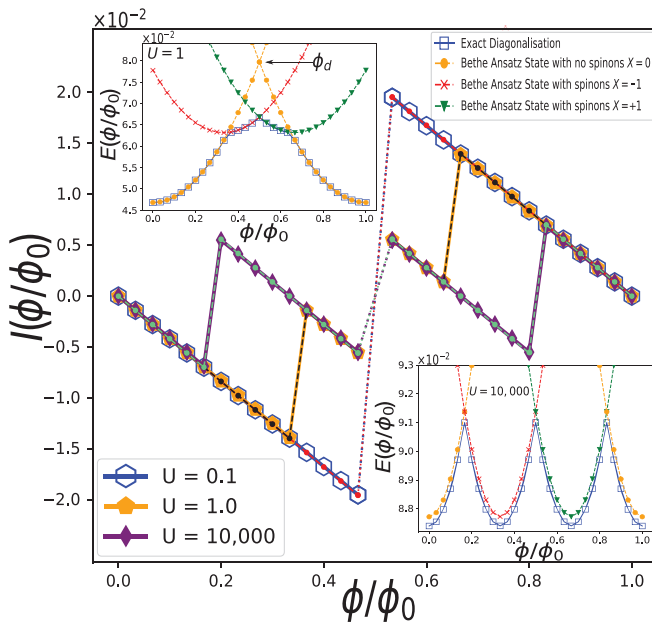
multicomponent  $SU(N)$  fermions is studied. The system is modeled by the  $SU(N)$  Hubbard model<sup>842</sup> with the repulsive interaction; the particles are confined in a ring-shape circuit pierced by an effective magnetic field. As discussed in Sec. VII, the zero temperature persistent current  $I(\phi)$  is defined as

$$I(\phi) = -\frac{\partial E_0}{\partial \phi}. \quad (139)$$

By applying a combination of Bethe ansatz<sup>842,855,856</sup> and numerical analysis, it is demonstrated how the persistent current displays a specific dependence on the parameters characterizing the physical conditions of the system. A combination of spin correlations, effective magnetic flux, and interaction brings about a peculiar phenomenon: spinon creation in the ground state.

The creation of spinons in the ground state leads to a redefinition of the elementary flux quantum  $\phi_0$ , which fixes the periodicity of the current. From Fig. 80, one can clearly observe how the profile of the persistent current changes with increasing  $U$ , which reflects the periodic  $1/N_p$  oscillations in the ground state energy that, in the large interaction regime, results in  $N_p$  parabolic cusps/segments.

Such fractionalization of the flux observed here is very different from the one typically observed in bosonic system with attractive interactions (see Sec. XIV): While the fractionalization in bosonic systems arises from the formation of bound states, for repulsing fermions, the phenomenon is a direct manifestation of the coupling between the spin and matter degrees of freedom.



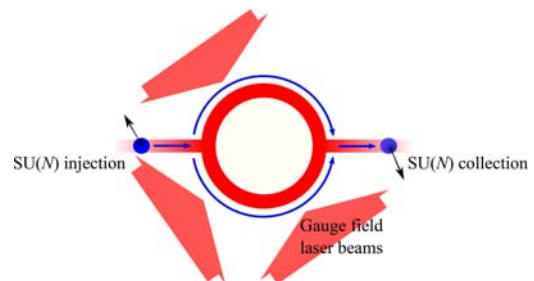
**FIG. 80.** Persistent current  $I(\phi)$  at incommensurate filling for  $SU(3)$  fermions with different interaction strengths  $U$  in the dilute filling regime of the Hubbard model. The exact diagonalization  $L = 30$ ,  $N_p = 3$  is monitored with the Bethe ansatz of the Sutherland–Gaudin–Yang model. The insets show how the Bethe ansatz energies need to be characterized by spinon quantum numbers in order to be the actual ground state. At  $U = 0$ , the ground state energy is a periodic sequence of parabolas meeting at degeneracy points  $\phi_d$  ( $\phi_d = 1/2$  for the case displayed in the figure). Figure is taken from Ref. 854.

Spinon creation in the ground state displays a marked dependence on the number of spin components, highlighting and distinguishing features between  $SU(2)$  and  $SU(N)$  fermions for  $N > 2$ . In particular, for integer fillings, at variance with their standard two spin component fermions counterpart,<sup>857</sup>  $SU(N)$  fermions with  $N > 2$  undergo a Mott quantum phase transition for a finite value of the interaction. Despite its mesoscopic nature, the persistent current is able to detect the onset of the Mott transition marked by a clear finite size scaling. Furthermore, the presence of a Mott gap suppresses spinon creation in the ground state. Finally, a specific  $SU(N)$  parity effect is shown to hold whereby the current is diamagnetic (paramagnetic) in nature for systems comprised of  $(2n + 1)N$   $[(2n)N]$  number of fermions, with  $n$  being an integer. This result generalizes a prediction by Byers–Yang, Onsager, and Leggett.<sup>811–813</sup>

## D. Concluding remarks and outlook

In Sec. XV A, we saw how to cool down alkaline-earth-like metal atoms to degeneracy and how we can take advantage of the fermionic isotopes and clock transition to address  $SU(N)$  Hamiltonian and engineer the light shifts between two orbital states almost at will. In Sec. XV B, we discussed the laser tripod scheme to implement (homogeneous) Abelian and non-Abelian gauge fields in the bulk [see Fig. 79(b)]. In this section, we discuss several potential research objectives and applications in elementary atomtronics circuit in the presence of  $SU(N)$  symmetry and/or gauge fields.

In the simplest instance, one can consider rectilinear and ring-shaped quasi-1D guides. The practical implementation can be done following methods discussed in Sec. II. Then, one can generate a non-Abelian synthetic gauge field in which the atomic spin will also change its orientation. This way, spin Hall current might be present even if the gauge field is constant and uniform in space<sup>88,851,858</sup> (see also Sec. XV B). This system can be implemented with various gauge structures, for example, uniform Abelian gauge field, spin–orbit configurations (in uniform non-Abelian gauge field), and synthetic magnetic fields such as a uniform or a monopole configuration. All these configurations can be realized using the tripod scheme developed for  $SU(2)$  systems. A natural extension of the tripod scheme can also be used for exploring the  $SU(3)$  symmetry.<sup>850</sup> One can fabricate non-Abelian Aharonov–Bohm matter-wave interferometers operating with a  $SU(N)$  fermionic fluid. To this end, one shall attach source and drain leads to the ring-shaped optical potential to inject and collect the quantum gas (see Fig. 81). Alternatively, the wires can be suppressed



**FIG. 81.** Sketch of a (non-Abelian)  $SU(N)$  Aharonov–Bohm matter-wave device. The tripod laser beams illuminate the full ring structure. With a single tripod scheme, we have  $N = 2$ . With a double tripod scheme, we have  $N = 3$ .

and the out-of-equilibrium dynamic of the ultracold gas can be investigated in the ring only. While the bosonic case has been largely studied (see Secs. III, VII, and XII and reference therein), the interacting fermionic case and the role of its spin internal structure remain largely unexplored. A number of theoretical questions need to be tackled to understand the dynamic of the system, for example, effects of the quasi-1D geometry, the interplay between charge and spin degrees of freedom at mesoscopic scales, the role of finite temperatures, and the impact of quantum statistics (Pauli blocking). Interestingly, persistent current states shall exist if pairing and superfluidity can occur and becomes superfluid and reaches the antiferromagnetic regime at strong repulsive interactions (Tonks regime). In the out-of-equilibrium regime, one can study the response to a quench in an isolated system. The integrable regimes could be explored by employing the machinery discussed in Secs. V and VI. One can also add localized barriers interrupting the ring (see Secs. III, VIII, IX, and X). Such a scheme would provide the implementation of an AQUID operating with a fermionic quantum fluid under a non-Abelian synthetic gauge field.

Based on the fermionic nature of the alkaline-earth atoms, it would be interesting to transpose the standard electronic and/or spintronics circuits to atomtronics circuits operating with neutral-atom fermionic species with enhanced control and flexibility.

The *primum mobile* for circuits with flowing  $SU(N)$  matter was theoretically analyzed recently.<sup>854</sup> It would be interesting to study configurations for atomic SQUIDs exploiting the  $SU(N)$  features. For  $SU(N)$  atomtronics, it would be also interesting to generalize the Datta-Das transistor (DDT), the fundamental building block of spintronics circuits,<sup>859</sup> to ultracold gas system. Major steps, toward that goal, were done theoretically,<sup>860</sup> and experimentally, on Rb BEC<sup>851</sup> and recently on strontium ultracold gas using a tripod scheme.<sup>861</sup> Among the possible specific added values of the latter implementation, a fermionic atomic DDT operating with an ultracold alkaline-earth-like gas can be extended to the gauge field with higher symmetry [ $SU(3)$  at least], which can be generated using a double tripod laser scheme.<sup>850</sup>  $SU(N)$  fermionic systems have triggered a great interest to explore their magnetic properties both theoretically<sup>840</sup> and experimentally.<sup>845</sup> One can exploit atomtronics circuits to probe  $SU(N)$  matter. For example, in the spirit of solid-state physics I-V characteristics, one could define a new route for the diagnostic of the different many-body quantum regimes in terms of the current flowing through the  $SU(N)$  system. Specifically, one could focus on fermionic systems that realize the  $SU(N)$  Heisenberg or Hubbard models in the rectilinear/ring-shaped potentials attached to source and drain leads. In these structures, the transport coefficients can be derived by monitoring the densities in the source and drain leads. One can consider investigating transport in the  $SU(N)$  Kondo impurity model. The effect of disorder, Anderson localization, and many-body localization<sup>862–865</sup> could be explored with fermionic atomtronics using  $SU(2)$  and  $SU(3)$  spin-orbit coupling. A similar logic could be employed to study the BCS-BEC crossover.<sup>866</sup>

## XVI. MANIPULATING RYDBERG ATOMS

W. Li, O. Morsch

Atoms excited to high-lying energy states (with principal quantum number  $n$  larger than  $\approx 15$ ) are known as Rydberg atoms.<sup>867</sup> They have considerably longer lifetimes than atoms in low-lying excited

states, and much larger (by several orders of magnitude, with strong scaling with  $n$ ) electric polarizability as well as dipole and van der Waals interactions. Rydberg atoms have been studied for several decades with renewed interest sparked by the invention of laser cooling, which made more accurate studies possible, and also due to the advent of quantum computation and quantum simulation, for which Rydberg atoms are a promising building block.<sup>572</sup> Generally, the combination of controllability, strong interactions and long coherence times make Rydberg atoms promising candidates for the realization of future quantum information technologies. In a broader context of quantum technologies, Rydberg atoms have also been explored for sensitive detection of electric fields and toward quantum state transfer between microwave and optical domains.

For the purposes of atomtronics, Rydberg atoms are an interesting system to study in regard to the propagation of excitations in disordered or ordered arrays. In fact, many transport properties in both the quantum and semiclassical regimes can be studied using Rydberg excitations. While Rydberg atoms have not been used for atomtronics applications (as understood in this review) so far, they might represent a valuable addition to the atomtronics toolbox in the future. In this spirit, the present section presents a few recent results on percolation phenomena studied in a gas of ultracold Rydberg atoms as well as on microwave control of Rydberg atoms.

### A. Driven-dissipative Rydberg systems

An important aspect of transport phenomena is the interplay between an external drive and the natural dissipation of the system, which has been investigated by several groups in recent years.<sup>868–872</sup> In samples of ultracold Rydberg atoms (with temperatures around  $T \approx 120 \mu\text{K}$ , so on the timescales of typical experiments atomic motion can be neglected), we can study this interplay by driving a transition between the ground state of the atom (87-rubidium in our case) and a high-lying Rydberg state with  $n \approx 70\text{--}80$ . In our experiments, in Pisa, we use  $S$  states (zero angular momentum) for which the van der Waals interaction is repulsive. This interaction leads to two distinct many-body effects. For resonant driving, it prevents the excitation of more than one Rydberg atom inside the “blockade sphere”; this is known as the dipole blockade.<sup>873,874</sup> On the other hand, for off-resonant driving, the van der Waals interaction can lead to the compensation of the detuning if a ground state atom is at a certain “facilitation distance” from a Rydberg atom.<sup>875,876</sup> At that distance, the off-resonant driving is shifted into resonance and, thus, the excitation of the ground state atom is “facilitated.”

It turns out that by adding the natural decay of a Rydberg state due to spontaneous emission (with timescales of a few hundred  $\mu\text{s}$ ), it is possible to realize a paradigmatic model from statistical physics called directed percolation,<sup>877</sup> which can be used to study such diverse processes as epidemic spreading, wildfires, or the onset of turbulence. This model can be characterized by two processes in a spin-1/2 example: offspring production in which a “spin up” causes a nearby “spin down” to flip its state at a certain rate and sudden death in which a spin up spontaneously flips down. For our Rydberg system, these two processes can be directly translated into facilitation with rate  $\Gamma_{\text{fac}}$  and spontaneous decay with rate  $\Gamma_{\text{spon}}$ . We note here that both processes are incoherent (in particular, we choose a Rabi frequency for Rydberg excitation that is smaller than the decoherence rate). From statistical physics, we know that this directed percolation model exhibits a phase



transition between its absorbing state (all spins “down,” or all atoms in the ground state) and an active state in which, on average, a macroscopic number of spins are “up” (i.e., atoms are in Rydberg states).

We realized this model using Rb Rydberg atoms in a magneto-optical trap.<sup>878</sup> By varying the Rabi frequency of the off-resonant laser driving (a two-photon excitation via an intermediate  $6P$  state was used), we were able to scan the ratio  $\Gamma_{\text{fac}}/\Gamma_{\text{spon}}$  across the critical value for the absorbing-state phase transition [ $\Gamma_{\text{fac}}$  is related to  $\Omega$  via  $\Gamma_{\text{fac}} = (\Omega^2/2\gamma)$ , where  $\gamma$  is the decoherence rate]. Figure 82 shows the results of those experiments. In order to prepare the system away from the absorbing state with all atoms in the ground state (from which, by definition, the system cannot escape), we initially excited around 30 Rydberg atoms in the cloud and then allowed the system to evolve under constant driving for 1.5 ms before measuring the number of Rydberg excitations by field ionization. The directed percolation phase transition is visible in both the plot of the number of excitations as a function of  $\Omega$  [Fig. 82(a)] and as a peak in the variance of the number of excitations [Fig. 82(b)].

This is one example of a transport/percolation problem implemented using cold Rydberg atoms. In future experiments, this concept can be extended to (partially) coherent driving<sup>879,880</sup> and/or ordered arrays<sup>881,882</sup> as well to tailored and controllable dissipation.

## B. Microwave-optical conversion using Rydberg atoms

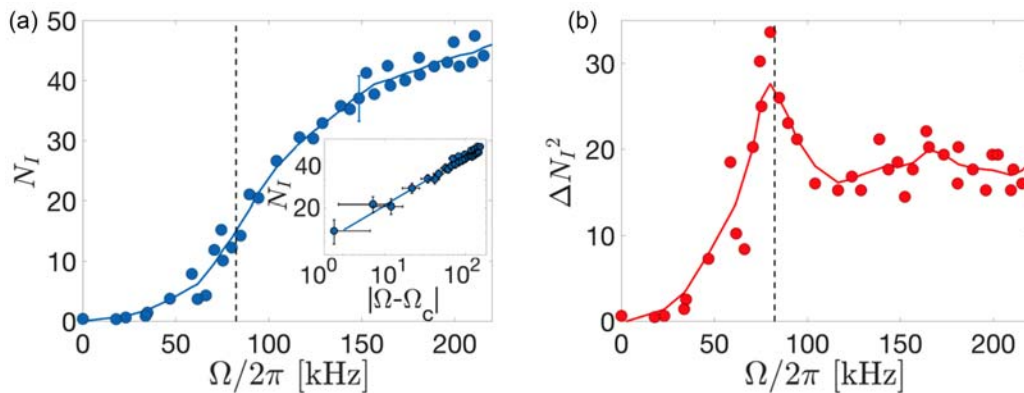
Rydberg atoms feature transitions of very large dipole moments in the microwave frequency range,<sup>867</sup> which has been utilized for sensitive detection of microwave electric field<sup>883,884</sup> and for efficient conversion from microwave to optical photons.<sup>885</sup> In quantum simulation using Rydberg atoms, nearby Rydberg states are commonly encoded as spin states, and their populations and dynamics can be conveniently manipulated with microwave radiation.<sup>27,886</sup>

Here, we present a demonstration of coherent microwave-to-optical conversion of classical fields via six-wave mixing in Rydberg atoms. In the quantum regime, such coherent conversion is essential for coupling superconducting qubits operating at microwave frequencies to photonic qubits used in quantum communication over long distances<sup>887</sup> and, therefore, has been intensively pursued in quite a few different physical systems.<sup>888</sup>

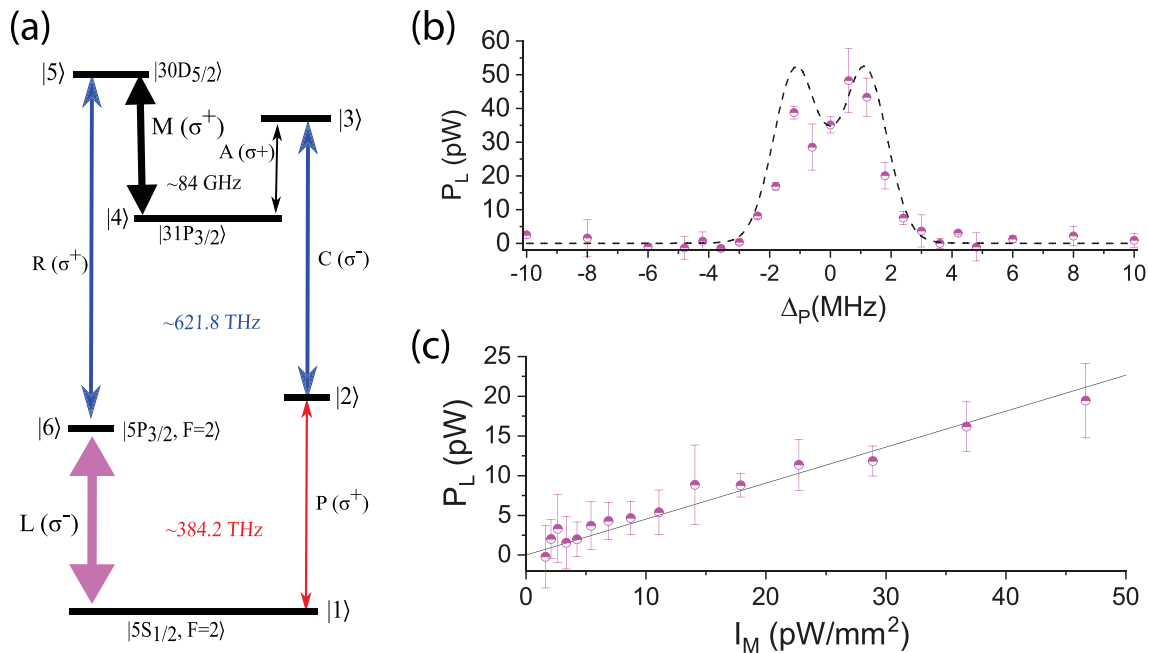
The principle of our conversion experiment using Rydberg atoms is as follows. A cloud of cold polarized <sup>87</sup>Rb atoms is illuminated by four auxiliary electromagnetic fields P, C, A, and R as well as the microwave field M to be converted. By nonlinear frequency mixing of the six waves in the atomic medium, the field M is converted into the optical field L. The chosen configuration of energy levels is displayed in Fig. 83(a), where the six waves are near-resonant with the atomic transitions shown in the figure with  $|1\rangle \equiv |5S_{1/2}, F=2, m_F=2\rangle$ ,  $|2\rangle \equiv |5P_{3/2}, F=3, m_F=3\rangle$ ,  $|3\rangle \equiv |30D_{3/2}, m_J=1/2\rangle$ ,  $|4\rangle \equiv |31P_{3/2}, m_J=-1/2\rangle$ ,  $|5\rangle \equiv |30D_{5/2}, m_J=1/2\rangle$ , and  $|6\rangle \equiv |5P_{3/2}, F=2, m_F=1\rangle$ . In the absence of the microwave field M, the system is in the configuration of microwave dressed electromagnetically induced transparency involving Rydberg states (Rydberg EIT), formed by the two optical waves P and C, and the auxiliary microwave field A. Once the M and R fields are added, the coherence induced between the ground state  $|1\rangle$  and the intermediate state  $|6\rangle$  triggers the generation of the converted optical field L.

A typical spectrum of the measured power  $P_L$  of the generated L field versus the input P field detuning  $\Delta_P$  is shown in Fig. 83(b). The conversion is most efficient around  $\Delta_P = 0$ , which is consistent with the nonlinearity responsible for the frequency mixing being maximum close to resonance. The behavior  $P_L$  for  $\Delta_P = 0$  is approximately linear as a function of the input intensity  $I_M$  of field M, as shown in Fig. 83(c). Given  $P_L \approx \alpha I_M$ , a linear fit to the data yields the photon conversion efficiency of the process to be  $\eta = 0.051$ . This conversion efficiency is 17 times larger than the one reported in Ref. 885, and this enhancement is due to an improved experimental configuration, which makes the conversion occur over a longer distance. Our theoretical study shows that by using a carefully selected energy level scheme to minimize the absorption of the input P field when propagating through the conversion medium, a conversion efficiency above 50% can be reached even with all-resonance six-wave mixing similar to that in Fig. 83(a).<sup>889</sup>

This conversion method is an application example from the strong coupling between microwave and Rydberg atoms. To reach near-unit conversion efficiency for quantum state transfer at the single photon level, one may consider implementing stimulated Raman adiabatic passage<sup>890</sup> or tuning two of the fields (for example fields C and A)



**FIG. 82.** Evidence for an absorbing state phase transition in a Rydberg gas. (a) Number of excitations in the stationary state as a function of  $\Omega$  (the solid line is a sliding average to guide the eye). The inset shows a power-law fit around the critical value  $\Omega_c$ , indicated by the dashed line in the main figure. In (b), the peak in the variance plotted as a function of  $\Omega$  indicates the critical point. Adapted with permission from Gutiérrez *et al.*, Phys. Rev. A **96**, 041602 (2017). Copyright 2017, American Physical Society.



**FIG. 83.** Efficient microwave-to-optical conversion using Rydberg atoms. (a) Energy level diagram and coupled transitions. The polarization of the fields are indicated inside parentheses. (b) Spectrum of the generated light power  $P_L$ . The dashed line is a simulated result obtained using Maxwell-Bloch equations. (c) The power of the generated light is plotted versus  $I_M$  in the range of 0 to 50 pW/mm<sup>2</sup>. The solid line is the result of a linear fit. Adapted with permission from Vogt *et al.*, Phys. Rev. A **99**, 023832 (2019). Copyright 2019, American Physical Society.

off-resonance to realize an effective two-photon transition in our system.<sup>891,892</sup> In addition to its potential for quantum state transfer, this method of conversion into optical photons is also promising for the sensitive real-time detection of microwave or THz fields.

### C. Concluding remarks and outlook

While the experiments outlined in this section do not yet make a direct contribution to atomtronics, it is likely that future studies of transport phenomena could make use of the techniques presented here. In the case of Rydberg atoms, Rydberg excitations coupled via dipole-dipole and van der Waals interactions—rather than the atoms themselves—are transported. Seed or source excitations can be injected into a cloud (or ordered array) of atoms at well-defined positions. In particular, using the recently developed patterning techniques based on dipole trap arrays (either using micromirror devices, holographic methods<sup>23</sup> or custom-made microlens-arrays<sup>154</sup>), it will be possible to conduct excitation-transport experiments using source-drain configurations (exploiting the high spatial resolution for excitation and detection). In this way, ring-shaped circuits or other more complicated transport topologies could be explored. Coupling of the Rydberg atoms to laser or microwave sources could then be used to further tailor the interaction between the Rydberg atoms or for inducing additional dissipation/dephasing in the system. This will allow one to study the crossover between incoherent hopping and coherent transport. Finally, it is conceivable that the studies of Rydberg excitation transport could be combined with “regular” atomtronics, resulting in a hybrid system in which both excitations and matter are transported (either independently or possibly coupled to each other).

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### DATA AVAILABILITY

Data sharing is not applicable to this article as no new data were created or analyzed in this study.

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